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(54) Title: RECOMBINANT CELL LINE

(57) Abstract: The invention relates to a recombinant cell line capable of inducible expression of an  $\alpha$  and/or  $\beta$  subunit of interleukin 12 (IL-12), and an ecdosymeinducible expression vector capable of transfecting a host cell to produce the recombinant cell line of the invention. The invention also relates to a method of screening a candidate compound for the ability to inhibit IL-12 formation and secretion which comprises the steps of incubating a cell line according to the invention with the candidate compound and then assaying the cell line culture for secreted IL-12, or a subunit thereof.



#### RECOMBINANT CELL LINE

#### Introduction

1	The	invention	also	relates	to	recombinant	cell	lines
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- 2 transformed to express a dimeric form of
- 3 interleukin, or a subunit thereof, and expression
- 4 vectors used to transform the cell lines. The
- 5 invention also relates to a method of screening
- 6 candidate compounds for the ability to inhibit
- 7 assembly and secretion of dimeric forms of
- 8 interleukins, or subunits thereof.

9

10

#### Background Art

.11

- 12 Cytokines are a unique family of growth factors.
- 13 Secreted primarily from leukocytes, cytokines
- 14 stimulate both the humoral and cellular immune
- 15 responses, as well as the activation of phagocytic
- 16 cells. Cytokines secreted from lymphocytes are
- 17 termed lymphokines, whereas those secreted by
- 18 monocytes or macrophages are termed monokines. Many
- 19 of the lymphokines are also known as interleukins

- 1 (IL's), since they are not only secreted by
- 2 leukocytes, but are also able to affect the cellular
- 3 responses of leukocytes. Specifically, interleukins
- 4 are growth factors targeted to cells of
- 5 hematopoietic origin. One of the interleukins, IL-
- 6 12, is a pro-inflammatory cytokine interleukin. This
- 7 cytokine is predominantly secreted either as a  $\alpha\beta$
- 8 heterodimeric form or as a  $\beta\beta$  homodimeric form. Both
- 9 dimer forms bind the IL-12-receptor on target cells
- 10 but differ in the spectrum of biological activities
- 11 induced. The  $\alpha\beta$  form is crucial for generation of
- 12 cell-mediated immunity against parasites, viruses
- 13 and bacteria, but contributes also to destructive
- 14 effects in pathogenesis of autoimmune diseases, e.g.
- 15 MS, RA and inflammatory bowel disease. The  $\beta\beta$  form
- 16 has been shown to be instrumental in virus-induced
- 17 inflammation, and in excessive epithelial airway
- 18 inflammation seen in asthma. Thus, both forms of IL-
- 19 12 are disease-promoting factors in a variety of
- 20 conditions. Recently, two novel cytokines have been
- 21 discovered, named interleukin-23 and interleukin-27
- 22 that apparantly belong to the IL-12 subclass of
- 23 cytokines based on structural relationships. Both
- 24 IL-23 and IL-27 share with IL-12 a typical
- 25 heterodimeric structure and are invloved in a
- 26 similar array of immune responses.

- 28 Celebrex is a diaryl-substituted pyrazole. It is a
- 29 nonsteroidal anti-inflammatory drug (NSAID) that is
- 30 indicated for the treatment of osteoarthritis,
- 31 rheumatoid arthritis, for the management of acute
- 32 pain in adults for the treatment of primary

- 1 dysmenorrhea. The mechanism of action of CELEBREX is
- 2 believed to be due to inhibition of prostaglandin
- 3 synthesis, primarily via inhibition of
- 4 cyclooxgenase-2 (COX-2). Scientific literature
- 5 indicates that CELEBREX displays antitumor effects
- 6 by sensitizing cancer cells to apoptosis. A recent
- 7 paper has indicated that CELEBREX blocks the
- 8 endoplasmic reticulum (ER) Ca2+-ATPases, and it has
- 9 been suggested that this Ca2+ perturbation may be
- 10 part of the signaling mechanism by which CELEBREX
- 11 triggers apoptosis. This Ca2+ perturbation effect
- 12 seems to be unique to CELEBREX and was not seen with
- 13 any of the other COX inhibitors (e.g. aspirin,
- 14 ibuprofen, naproxen etc.)

16 Statement of Invention

17

- 18 According to the invention, there is provided an
- 19 expression vector comprising DNA encoding a subunit
- 20 of a dimeric form of interleukin under
- 21 transcriptional control of an ecdysone-inducible
- 22 promoter.

23

- 24 Suitably, the subunit of a dimeric form of
- 25 interleukin is selected from the group comprising:
- 26 p35 (alpha) subunit of interleukin 12 (IL-12); p40
- 27 (beta) subunit of IL-12; p19 chain of IL-23; p40
- 28 subunit of IL-23; ebi3 subunit of IL-27; and p28
- 29 subunit of Il-27.

- 31 Typically, the vector comprises an ecdysone-
- 32 inducible mammalian expression plasmid, wherein the

DNA encoding the subunit of a dimeric form of interleukin is included in the plasmid.

3

- 4 In one embodiment of the invention, the vector
- 5 comprises DNA encoding a p40 subunit of IL-12. Cell
- 6 lines stably transfected with such a vector will,
- 7 when induced, express both homodimeric IL-12 and the
- 8 beta-subunit of IL-12.

9

- 10 In another embodiment of the invention, the vector
- 11 comprises DNA encoding a p35 subunit of IL-12. Cell
- 12 lines stably transfected with such a vector will,
- 13 when induced, express the alpha-subunit of IL-12.

14

- 15 In another embodiment of the invention, the vector
- 16 comprises DNA encoding a p19 subunit of IL-23. Cell
- 17 lines stably transfected with such a vector will,
- 18 when induced, express the p19 subunit of IL-23.

19

- 20 In a preferred embodiment of the invention, the
- 21 ecdysone inducible mammalian expression vector is
- 22 selected from the group comprising: pIND; pIND(SP1);
- 23 and pINDHygro.

24

- 25 In a particularly preferred embodiment of the
- 26 invention, the DNA encoding a subunit of dimeric
- 27 interleukin 12 includes a DNA sequence encoding a 6
- 28 x histidine tag.

- 30 In one embodiment of the invention, the expression
- 31 vector is selected from the group comprising: pIND-

1 p35H; pIND(SP1)-p35H; pIND-40H; pINDHygro-p40; 2 pIND(SP1)-p40H; and pIND-p40. 3 Suitably, the DNA encoding the subunit of dimeric 4 interleukin is digested with NheI and XhoI 5 restriction enzymes prior to ligation of the 6 digested DNA products into the expression vector. 7 8 The invention also relates to an expression vector 9 pIND(SP1)-p35H having ECACC accession number 10 03120401. A sample of this vector was deposited at 11 the ECACC on 4 December 2003. 12 13 The invention also relates to a method a producing a 14 tightly controlled expression vector capable of 15 transforming a host cell which when transformed is 16 capable of producing a recombinant dimeric 17 18 interleukin, or a subunit thereof, under transcriptional control of a ecdosone inducible 19 20 promoter, comprising the steps of: 21 - providing cDNA for a subunits of a dimeric 22 interleukin; 23 - digesting the cDNA with at least one 24 restriction enzyme; and 25 - ligating the digested cDNA product into an ecdysone-inducible mammalian expression vector. 26 27 In a preferred embodiment of the invention, the DNA 28 is digested with two restriction enzymes, these 29 being NheI and XhoI. Suitably, the plasmid into 30 which the digested DNA is to be ligated is also 31

digested with the same restriction enzymes.

- 2 The invention also relates to an expression vector
- 3 obtainable by the method of the invention.

4

- 5 The invention also relates to a cell line
- 6 transfected with at least one expression vector of
- 7 the invention, wherein the DNA encoding the at least
- 8 one subunit of a dimeric interleukin is under
- 9 transcriptional control of a ecdysone-inducible
- 10 mammalian expression system.

11

- 12 Suitably, the ecdysone-inducible mammalian
- 13 expression system comprises a plasmid other the
- 14 expression vector of the invention which
- 15 constitutively expresses two receptors which
- 16 interact in the presence of ecdysone, or an analog
- 17 thereof, to form a complex which binds to a response
- 18 element of a promotor controlling DNA encoding the
- 19 at least one subunit of a dimeric interleukin. Such
- 20 a plasmid is sold by Invitrogen under the name
- 21 pVgRxR.

22

- 23 In one embodiment, the cell line is transfected with
- 24 DNA that encodes a p35 (beta) subunit of IL-12. Such
- 25 a cell line, when induced, produces homodimeric IL-
- 26 12 and the beta-subunit of IL-12.

- 28 In another embodiment, the cell line is transfected
- 29 with an expression vector which includes DNA
- 30 encoding the p40 subunit of IL-12, and a further
- 31 expression vector which includes DNA encoding the

- 1 p35 subunit of IL-12. Such a cell line, when
- 2 induced, produces heterodimeric IL-12.

3

- 4 In another embodiment, the cell line is transfected
- 5 with an expression vector which includes DNA
- 6 encoding the p40 subunit of IL-12 (which is
- 7 identical to the p40 subunit of IL-23), and a
- 8 further expression vector which includes DNA
- 9 encoding the p19 subunit of IL-23. Such a cell line,
- 10 when induced, produces heterodimeric IL-23.

11

- 12 Typically, the cell lines of the invention include
- 13 the plasmid pVgRxR.

14

- 15 In one embodiment of the invention, the cells of the
- 16 cell line are human embryonic kidney cells,
- 17 preferably EcR293 cells.

18

- 19 The invention also relates to a cell line according
- 20 to the invention in which the cells are natural
- 21 beta-subunit-producing cells such as a HIBERNIA1
- 22 cell line.

23

- 24 The invention also relates to a cell line having
- 25 ECACC accession number 03112701. This cell line
- 26 includes an expression vector having DNA encoding
- 27 for the p40 (beta) subunit of IL-12. A deposit of
- 28 the recombinant cells was made at the ECACC on 27
- 29 November 2003.

- 31 The invention also relates to a method of producing
- 32 a cell line capable of producing a recombinant

1	dimeric interleukin, or a subunit thereof, under
2	transcriptional control of a ecdysone-inducible
3	promoter, comprising the steps of:
4	<ul> <li>providing at least one expression vector</li> </ul>
5	according to the invention; and
6	- transfecting a host cell with the at least one
7	expression vector,
8	wherein the DNA encoding the at least one subunit
9	of a dimeric interleukin is under the
10	transcriptional control of a ecdysone-inducible
11	mammalian expression system.
12	
13	The invention also relates to a method of preparing
14	cDNA encoding a subunit of a dimeric form of
15	interleukin comprising the steps of providing cDNA
16	encoding the subunit, and digesting the cDNA with
17	restriction enzymes NheI and XhoI to obtain a cDNA
18	product.
19	
20	The invention also relates to a method of screening
21	a candidate compound for the ability to inhibit
22	dimer assembly and secretion of a dimeric form of
23	interleukin, comprising the steps of:
24	<ul> <li>incubating a cell culture comprising a cell</li> </ul>
25	line of the invention with the candidate
26	compound;
27	<ul> <li>inducing transcription of the dimeric</li> </ul>
28	interleukin in the cells of the culture using

ecdysone or an ecdysone analog; and

- assaying the cell culture for the presence of
secreted interleukin.

- 1 In one embodiment of the method, the interleukin
- 2 expressed by the cell line has a 6 x histidine amino
- 3 acid sequence tagged on either or both of the
- 4 subunits thereof, wherein the assaying step involves
- 5 Ni-NTA affinity chromatography.
- 6 Alternatively, the assaying step involves probing
- 7 the cell culture with an antibody specific to a
- 8 dimeric form of interleukin, or a subunit thereof.

- 10 The invention also relates to an inhibitor of dimer
- 11 assembly and secretion of dimeric interleukin
- 12 identified by the method of the invention.

13

- 14 The invention also relates to a method of prevention
- 15 or treatment of inflammatory disease comprising a
- 16 step of treating an individual with an inhibitor
- 17 identified by the method of the invention. One such
- 18 inhibitor IDENTIFIED is CELEBREX.

19

- 20 In a further aspect, the invention provides a method
- 21 of treating disease having a pathogenesis which
- 22 includes endogenous production of any of cytokines
- 23 IL-12, IL 23 or IL-27, the method comprising a step
- 24 of treating an individual with an endoplasmic
- 25 reticulum (ER) Ca<sup>2+</sup> perturbation reagent.

26

- 27 In a further aspect, the invention provides the use
- 28 of an ER Ca2+ perturbation reagent in the manufacture
- 29 of a medicament for the treatment of disease having
- 30 a pathogenesis which includes endogenous production
- of any of cytokines IL-12, IL-23 or IL-27.

- 1 In a further aspect, the invention provides the use
- 2 of an ER Ca<sup>2+</sup> perturbation reagent for the treatment
- 3 of disease having a pathogenesis which includes
- 4 endogenous production of any of cytokines IL-12, IL-
- 5 23 or IL-27.

- 7 In a further aspect, the invention relates to a
- 8 method of inhibiting the formation of one or more
- 9 cytokines in an individual, which method comprises
- 10 the step of treating an individual with ER Ca2+
- 11 perturbation reagent. In one embodiment, the
- 12 cytokines are selected from IL-12, IL-23 and IL-27.

13

- 14 In a further aspect, the invention relates to the
- 15 use of an ER  $Ca^{2+}$  perturbation reagent to inhibit the
- 16 formation of one or more cytokines in an individual.
- 17 In one embodiment the cytokines are selected from
- 18 IL-12, IL-23 and IL-27.

19

- 20 In a preferred embodiment, the disease is an
- 21 inflammatory disease. More preferably, the disease
- 22 is a disease in which one or more endogenously
- 23 produced IL-12 forms play a disease promoting role.
- 24 Typically, the IL-12 forms are  $\alpha\beta$  heterodimeric and
- 25  $\beta\beta$  homodimeric forms.

26

- 27 In one embodiment, diseases in which cyclooxygenase-
- 28 2 (COX-2) is reported to play a substantial disease
- 29 promoting role are disclaimed.

- 31 In one embodiment, the inflammatory disease is a
- 32 disease in which the endogenous production of one or

- 1 both of  $\alpha\beta$  and  $\beta\beta$  forms of IL-12 is known to lead to
- 2 disease in a COX-2 independent manner.

3

- 4 The invention also relates to a method of inhibiting
- 5 the production of one or more cytokines in an
- 6 individual in a post-translational manner, which
- 7 method comprises a step of treating an individual
- 8 with ER Ca<sup>2+</sup> perturbation reagent.

9.

- 10 Preferably, the disease is selected from the group
- 11 consisting of infectious diseases; bacterial
- 12 protozoal or virus-induced inflammation; epithelial
- 13 airway inflammation such as asthma; allergic
- 14 disease; autoimmune disease such as MS, RA and
- 15 Inflammatory Bowel Disease; and -all conditions in
- 16 which endogenously produced IL-12  $\alpha/\beta$  or  $\beta\beta$  forms
- 17 are thought to play a disease-promoting role,
- 18 including:

- 20 Pulmonary fibrosis
- 21 Pulmonary tuberculosis
- 22 Asthma
- 23 Sarcoidosis
- 24 Leprosy
- 25 Schistosomiasis
- 26 Lupus erythematosis
- 27 Lupus nephritis
- 28 Allograft rejection
- 29 Airway inflammation
- 30 Respiratory syncytial virus infection
- 31 Multiple sclerosis
- 32 Alzheimer's disease

- 1 Abortion (women with recurrent pregnancy loss)
- 2 Certain vaccines aimed at inducing TH2-type immune
- 3 responses
- 4 Experimental autoimmune myocarditis
- 5 Tuberculosis
- 6 Psoriatic arthritis
- 7 Rheumatoid arthritis
- 8 Osteoarthritis
- 9 Colonic inflammation (colitis)
- 10 Crohn's Disease
- 11 Inflammatory bowel disease
- 12 Atopic dermatitis, AD (chronic stage)
- 13 Inflammatory skin disease
- 14 Insulin dependent diabetes mellitus Type I and II
- 15 Endotoxaemia
- 16 Exposure to organic dust
- 17 Periodontal diseases
- 18 Nephrotic syndrome
- 19 Hepatocellular damage in chronic hepatitis C
- 20 Primary biliary cirrhosis
- 21 Cancer patients (Various cancers, and various stages
- 22 in cancer that are typically accompanied with
- 23 dysregulated IL-12, IL-23 and/or or IL-27
- 24 production)
- 25 ANCA associated vasculitis and sepsis
- 26 Experimental crescentic glomerulonephritis
- 27 Atherosclerosis
- 28 Acute viral myocarditis
- 29 Autoimmune myocarditis
- 30 Experimental autoimmune myastenia gravis
- 31 Uveitis (as Behret's disease)
- 32 Thyroiditis and Grave's disease

- 1 Thyroid autoimmune disease
- 2 Myelopathy (HTLV-I-associated myelopathy)
- 3 Symptomatic transient hypogammaglobulinaemia of
- 4 infancy (THI)
- 5 Selective IgA deficiency (SIgAD)
- 6 Schizophrenia
- 7 Primary malignant melanoma
- 8 Abdominal aortic aneurysm
- 9 Autoimmune thrombocytopenic purpura
- 10 Heatstroke
- 11 Meningococcal sepsis
- 12 Septic shock
- 13 Meningoencephalitis
- 14 Bacterial meningitis
- 15 Pregnancy
- 16 Pre-eclampsia
- 17 HELLP syndrome (hemolysis, elevated liver function
- 18 test and low platelet counts
- 19 Endometriosis
- 20 Acute pancreatitis
- 21 Lung fibrosis induced by silica particles
- 22 Scleroderma
- 23 Sjogren's syndrome
- 24 Ankylosis spondylitis
- 25 Hashimoto's thyroiditis
- 26 Autimmune anemias
- 27 Goodpasture's syndrome
- 28 Addinson's disease
- 29 Autoimmune hemolitic anemia
- 30 Spontaneous infertility (sperm)
- 31 Poststreptococcal glomerulonephritis
- 32 Autoimmune neuritis (Guillian-Barrd syndrome)

- 1 Sialadenitis
- 2 Brucellosis
- 3 Chickenpox and related viral diseases
- 4 Helicobacter Pyloris-induced gastritis
- 5 Common Variable Immunodeficiency (CVI)

6

- 7 In one embodiment, the disease is a conditions
- 8 characterized by dysregulation of IL-12, IL-23 or
- 9 IL-27 production conferred by polymorphisms in their
- 10 respective genes, or by polymorphisms in genes
- 11 involved in the biological activation or signal
- 12 transduction pathway of these cytokines.

13

- 14 In one embodiment, the ER Ca<sup>2+</sup> perturbation reagent
- 15 is selected from the compounds of Formula I:

16

17 Formula I

$$R^2$$
 $S$ 
 $R^3$ 
 $R^3$ 

- 19 wherein A is a substituent selected from partially
- 20 unsaturated or unsaturated hetrocyclyl and partially
- 21 unsaturated or unsaturated carbocyclic rings;
- 22 wherein R<sup>1</sup> is at least one substituent selected from
- 23 hetercyclyl, cycloalkyl, cycloalkenyl and aryl,
- 24 wherein R<sup>1</sup> is optionally substituted at a
- 25 substitutable position with one or more radicals
- 26 selected from alkyl, haloalkyl, cyano, carboxyl,
- 27 alkoxycarbonyl, hydroxyl, hydroxyalkyl, amino,
- 28 alkylamino, arylamino, nitro, alkoxyalkyl,
- 29 alkylsulfinyl, halo, alkoxy and alkylthio;

- 1 wherein R<sup>2</sup> is methyl or amino; and
- 2 wherein  $R^3$  is a radical selected from hydrido, halo,
- 3 alkyl, alkenyl, oxo, cyano, carboxyl, cyanoalkyl,
- 4 heterocyclyloxy, alkyloxy, alkylthio, alkylcarbonyl,
- 5 cycloalkyl, aryl, haloalkyl, heterocyclyl,
- 6 cycloalkenyl, aralkyl, hetrocyclylalkyl, acyl,
- 7 alkythioalkyl, hydroxyalkyl, alkoxycarbonyl,
- 8 arylcarbonyl, aralkylcarbonyl, aralkenyl,
- 9 alkoxyalkyl, arylthioalky, aryloxyalkyl,
- 10 aralkylthioalky, aralkoxyalkyl, alkoxyaralkoxyalkyl,
- 11 alkoxycarbonalkyl, aminocarbonyl,
- 12 aminocarbonylalkyl, alkyaminocarbonyl, N-
- 13 arylaminocarbonyl, N-alkyl-N-arylaminocarbonyl,
- 14 alkylaminocarbonylalkyl, carboxyalkyl, alkylamino,
- 15 N-arylamino, N-aralkylamino, N-alkyl-N-aralkylamino,
- 16 N-alkyl-N-arylamino, aminoalkly, alkylaminoalkyl, N-
- 17 arylaminoalkyl, N-aralkylaminoalkyl, N-alkyl-N-
- 18 aralkylaminoalky, N-alkyl-N-arylaminoalkyl, aryloxy,
- 19 aralkoxy, arylthio, aralkylthio, alkylsulfinyl,
- 20 alkylsulfonyl, aminosulfonyl, alkylaminosulfonyl, N-
- 21 arylaminosulfonyl, arylsulfonyl, N-alkyl-N-
- 22 arylaminosulfonyl; or a pharmaceutically-acceptable
- 23 salt thereof.

- 25 In a preferred embodiment, the ER Ca<sup>2+</sup> perturbation
- 26 reagent is selected from the compounds and
- compositions described in US Patent 5,972,986,
- 28 Column 3, line 34 to Column 10, line 32. In a
- 29 particularly preferred embodiment, the ER Ca<sup>2+</sup>
- 30 perturbation reagent is a diaryl- substituted
- 31 pyrazole marketed under the brand name CELEBREX
- 32 (Celecoxib). CELEBREX is chemically designated as 4-

- 1 [5-(4-methylpheny)-3-(trifluoromethyl)-IH-pyrazol-I-
- 2 y1] benzenesulfonamide.

3

- 4 Alternatively, the ER Ca2+ perturbation reagent may
- 5 be thapsigargin or A23187.

6

- 7 The invention will be more clearly understood from
- 8 the following description of some embodiments
- 9 thereof, given by way of example only, with
- 10 reference to the accompanying figures.

11

12 Brief Description of the Figures

13

- 14 Figure 1. is a schematic representation of the
- 15 Ecdysone-Inducible Mammalian Expression System.

16

- 17 Figure 2. is a schematic overview of the pIND,
- 18 pINDSP1 and pINDHygro vectors.

19

- 20 Figure 3. Primers used for amplification of the  $\boldsymbol{\alpha}$
- 21 and  $\beta$  chains of IL-12. (A) $\alpha$  chain forward primer;
- 22 (B)lpha -chain reverse primer; (C)eta chain forward
- 23 primer; (D)  $\beta$ -chain reverse primer and (E)  $\beta$ -chain
- 24 reverse primer without histidine tag. The sequence
- 25 coding for the hexahistidine-tag is represented in
- 26 red, while initiation and stop codons are indicated
- 27 in bold. The Kozak translation initiation sequence
- 28 is underlined.

- 30 Figure 4. Analysis of the amplification of the  $\beta$ -
- 31 chain from LPS-induced U937 cells by means of 1.5%

- 1 agarose gel electrophoresis. Lane 1, 100-bp DNA
- 2 marker; Lane 2-4,  $\beta$ -chain fragment amplified in the
- 3 presence of 2 mM MgSO<sub>4</sub> (lane 2); 3 mM MgSO<sub>4</sub> (lane 3)
- 4 or 4 mM MgSO<sub>4</sub> (lane 4).

- 6 Figure 5. Amplification of  $\alpha$ -chain cDNA (702bp).
- 7 Lane 1, 100-bp DNA marker; Lane 2-4, α-chain
- 8 fragment amplified in the presence of Pwo DNA
- 9 polymerase and 2 mM MgSO<sub>4</sub> (lane 2); 3 mM MgSO<sub>4</sub> (lane
- 10 3) or 4 mM MgSO<sub>4</sub> (lane 4).

11

- 12 Figure 6. Amplification of  $\beta$ -chain cDNA (1029bp).
- 13 Lane 1, 100-bp DNA marker; Lane 2-4, β-chain
- 14 fragment amplified in the presence Pwo DNA
- 15 polymerase of 2 mM MgSO4 (lane 2); 3 mM MgSO4 (lane
- 16 3) or 4 mM MgSO<sub>4</sub> (lane 4). Lanes 1-3 correspond to
- 17 products obtained using the reverse primer without
- 18 the histidine tag and lanes 5-6 including the
- 19 histidine tag.

20

- 21 Figure 7. Expression cassettes for the  $\alpha$  and  $\beta$ -
- 22 chains of IL-12 in the series of pIND vectors. (A)
- 23 Expression cassette shared by all vectors of the
- 24 pIND series with indication of the location of the
- 25 minimal heat shock promoter  $(P_{\Delta HSP})$  and the bovine
- 26 growth hormone poly-adenylation signal (BGH pA); (B)
- 27 and (C) 5' and 3' nucleotide sequences and
- 28 corresponding amino- and carboxy-terminal amino acid
- 29 sequences of the recombinant  $\alpha$  (B) and  $\beta$  (C) chains
- 30 with indication of the primer sequences.

- 1 Figure 8. Electrophoresis of amplification products
- 2 obtained by colony PCR of ampicillin-resistant
- 3 clones. The photographs show the results obtained
- 4 from clones transformed with (A) pIND(SP1)-p40H; (B)
- 5 pINDHygro-p40; and (C) pIND-p40.

- 7 Figure 9. Electrophoresis of amplification products
- 8 obtained by colony PCR of ampicillin-resistant
- 9 clones following transformation with pIND(SP1)-p35H

10

- 11 Figure 10. Confirmation of the presence of inserts
- 12 by means of restriction analysis of minipreps. (M)
- 13 100-bp ladder; (A) pIND(SP1)-p35H digested with NheI
- 14 and XhoI (insert of 700 bp); (B) pINDHygro-p40
- 15 digested with NheI and XhoI; and (C) pIND(SP1)-p40H
- 16 digested with NheI and XhoI (inserts of 900 bp).
- 17 Note: the vector portions were too large to
- 18 penetrate into this high-percentage agarose gel and
- 19 are therefore not visible.

- 21 Figure 11. Analysis of ponasterone A-inducible
- 22 expression of IL-12 lpha (A) and eta (B) chains in
- 23 transfected cell lines. 4-15% reducing SDS-PAGE
- 24 analysis of clones 1A9 (His-tagged  $\alpha$  -chain), 2G10
- 25 (His-tagged  $\alpha$ -chain) and 3D9 ( $\beta$ -chain). (A)
- 26 detection with monoclonal anti-p35 antibody. 1 (lane
- 27 1), 5 (lane 2) and 10 (lane 3) µl of the medium, and
- 28 1 (lane 4), 5 (lane 5) and 10 µl (lane 6) of the
- 29 soluble cell lysate of ponasterone A-induced clone
- 30 1A9 were submitted to 4-15% SDS-PAGE and
- 31 immunoblotted. Lanes 7-12 represent similar
- 32 fractions of clone 2G10. (B) detection with

- 1 monoclonal anti-p40 antibody. Lanes 1-6: fractions
- 2 of medium and cell lysate of clone 3D9 as described
- 3 for (A); Lanes 7-12: cell lysates of clones 1A9 and
- 4 2G10, used as negative control.

- 6 Figure 12. Expression levels of the IL-12  $\alpha$  chain in
- 7 18 different neomycin-resistant EcR293 clones.
- 8 Anti- $\alpha$ -chain immunoblots of soluble cell lysates
- 9 were prepared from induced (I) and uninduced (U)
- 10 EcR293 clones obtained following transfection and
- 11 neomycin selection with (A) pIND-p35H; (B, C)
- 12 pIND(SP1)-p35H and (D) pIND-p35H or pIND(SP1)-p35H.
- 13 Lysates were subjected to reducing SDS-PAGE using 4-
- 14 15% gels, blotted and immunodetected with anti lpha-
- 15 chain antibody. As negative control, we used the
- 16 secreted fraction of clone 4B6Z, which expresses the
- 17  $\beta$ -chain (lane 13-14 in Figure 16D).

18

- 19 Figure 13. Expression levels of the IL-12  $\beta$  chain in
- 20 hygromycin- (A) and neomycin- (B) resistant EcR293
- 21 clones. Anti- $\beta$ -chain immunoblots of soluble cell
- 22 lysates prepared from induced (I) and uninduced (U)
- 23 EcR293 cells. Clones were obtained by transfection
- 24 with (A) pINDHygro-p40; or (B) pIND(SP1)-p40H.
- 25 Lysates were subjected to SDS-PAGE using 4-15% gels,
- 26 blotted and immunodetected with anti  $\alpha$ -chain
- 27 antibody.

- 29 Figure 14. Transient transfection of HIBERNIA.1
- 30 cells with pIND(SP1)-p35H. Non-reducing 4-15% SDS-
- 31 PAGE and immunoblot of secreted fractions of the

- 1 transfected cell line following 30 (lanes 1 and 2)
- 2 and 48 (lanes 3 and 4) hrs of induction with
- 3 ponasterone A. The cells were transfected with 1
- 4 (lanes 1 and 3) or 2 (lanes 2 and 4)  $\square g$  of
- 5 pIND(SP1)-p35H. As a control the secreted fraction
- 6 of the non-transfected induced  $\beta$ -chain-producing
- 7 HIBERNIA.1 cells was used (lane 5). (A) detection
- 8 with anti  $\beta$ -chain antibody; (B), detection with anti
- 9  $\alpha$ -chain antibody.

- 11 Figure 15. Immunodetection of lpha and eta subunits of
- 12 IL-12 in medium of HIBERNIA.1 cells transiently
- 13 transfected with pIND(SP1)-p35H following reducing
- 14 SDS-PAGE. Lane 1, detection with anti  $\alpha$ -chain
- 15 antibody; Lane 2, detection with anti- $\beta$ -chain
- 16 antibody, Lane 3, detection with both antibodies at
- 17 the same time.

18

19 Detailed Description of the Invention

- 21 Recombinant cell lines that secrete various forms of
- 22 IL-12 under control of tightly regulated promoters
- 23 were generated. It was observed that treatment of
- 24 these cell lines with an ER Ca2+ perturbation reagent
- 25 such as thapsigarin inhibited secretion of both the
- 26  $\alpha\beta$  and  $\beta\beta$  forms of IL-12. The compound CELEBREX was
- 27 also tested on assembly of IL-12, and found that it
- 28 exerts a similar inhibitory effect on the secretion
- 29 of the  $\alpha\beta$  and  $\beta\beta$  forms of IL-12. There is a total
- 30 block in the secretory production of both dimer
- 31 forms of IL-12, and maximal effects are obtained

- 1 with the normal physiological working concentration
- 2 of CELEBREX in the absence of any apparent toxic
- 3 effects as measured with the MTT assay. These
- 4 affects are conferred in a post-transcriptional and
- 5 post-translation manner as there is no effect on
- 6 mRNA of IL-12. Without being bound by theory,
- 7 evidence has been produced to support a Ca2+ -
- 8 dependent disturbance in the folding pathway of IL-
- 9 12 due to impaired activity of certain chaperones in
- 10 the ER.

- 12 The inhibitory effect of CELEBREX on formation of
- 13 the  $\alpha\beta$  and  $\beta\beta$  forms of IL-12 in vitro indicates that
- 14 this drug is of interest for the treatment of
- 15 inflammatory conditions in which endogenous
- 16 production of these IL-12 forms is known to lead to
- 17 disease in a COX2-independent manner, including MS,
- 18 IBD, virus-induced inflammation and asthma.

- 20 IL-12 is a member of a family of cytokines that
- 21 includes two recently discovered members IL-23 and
- 22 IL-27. All of these cytokines have a typical
- 23 heterodimeric structure and display an array of both
- 24 overlapping and distinct activities. It is thought
- 25 that also IL-23 and IL-27 may contribute to
- 26 destructive inflammation in various conditions.
- 27 Most anti-cytokine drugs work by inhibiting
- 28 transcription of mRNA. To our knowledge this is the
- 29 first demonstration of a drug that inhibits cytokine
- 30 formation in a post-translational manner on the
- 31 level of folding and secretion of the protein, i.e.
- 32 by perturbation.

1 2 Experimental methods 3 4 Materials. Celecoxib (Celebrex) was obtained from Hefei Sceneri Chemical Co.; thapsigargin was 5 obtained from Calbiochem and A23187 from Sigma. 6 7 Cell culture. HEK293 IL-12  $\beta/\beta$  and  $\alpha/\beta$  producing 8 cell lines were maintained in a CO2 incubator at 37 9 10  $^{\circ}$ C (5% CO<sub>2</sub>). Cells were cultured in DMEM medium supplemented with 10% foetal bovine serum. 11 12 Cloning and expression of the  $\alpha$  and  $\beta$  chain of IL-12 13 14 Extraction of mRNA from IL-12 producer cell line ' 15 16 Human monocytic U937 cells were kindly provided by 17 the Rega Institute, Leuven, Belgium. U937 cells were 18 grown in DMEM (Dulbecco's modified eagle medium) 19 20 supplemented with 10응 FBS, 2 mM L-glutamine 21 (LifeTechnologies) and 50 µg/ml of gentamycin (Sigma). Cells were cultivated in 75cm<sup>2</sup> flasks, in a 22  $CO_2$  incubator (5%  $CO_2$ ) at 37°C and subcultured once a 23 week by splitting 1/10 by means of trypsination with 24 25 Trypsin-EDTA (LifeTechnologies) followed centrifugation to remove trypsin. Cells were induced 26 with IFN-  $\gamma$  (100 ng/ml) and LPS (1  $\mu$ g/ml; Sigma) for 27 24 hours. Total RNA was extracted from cells  $(10^7)$ 28 29 using StrataPrp® Total RNA Miniprep kit 30 (Stratagene). This method uses а powerful denaturant, guanidine thiocyanate, 31 in the lysis buffer. Afterwards, the sample was filtrated to 32

- 1 reduce the amount of DNA and subjected to a silica-
- 2 based fibre matrix to capture RNA.

4 Amplification of  $\alpha$  and  $\beta$ -chains of IL-12 by RT-PCR

5

- 6 To perform RT-PCR on the RNA extracted from IL-12
- 7 producer cells, we used the ProSTAR™ HF Single-Tube
- 8 RT-PCR System (High Fidelity) obtained from
- 9 Stratagene. This method uses the StrataScript
- 10 reverse transcriptase, which is subsequently
- 11 inhibited by incubation at 95°C. Amplification is
- 12 achieved with TaqPlus Precision polymerase.
- 13 Oligonucleotides complementary to the sequences to
- 14 be amplified ( $\alpha$  and  $\beta$ -chain) were synthesized by
- 15 LifeTechnologies. For the  $\alpha$ -chain, the forward
- 16 primer was designed to contain the second initiation
- 17 methionine (ATG) and NheI restriction site (GCTAGC),
- 18 while the reverse primer contained the stop codon
- 19 (TAA), XhoI restriction site (CTCGAG) and a 6 imes
- 20 Histidine tag sequence [3x(ATGGTG) ]. The  $\beta$ -chain
- 21 forward primer contained the initiation codon and
- 22 the NheI restriction site as well. We synthesized
- 23 two different oligonucleotides as reverse primers.
- 24 The first one contains the stop codon, XhoI
- 25 restriction site and the 6xHis sequence, and the
- 26 second was designed without the 6×Histidine
- 27 sequence.

28

29 α-chain

- 31 Forward 5'CAGGCTAGCGCAGCCATGTGTCCAGCGCGCAGC3'
- 32 Reverse 5'CTGCTCGAGTTAATGGTGATGGTGATGGTGGGAAGCA

1	TTCAGATAGCT3'						
2	$\beta$ -chain						
3							
4	Forward 5'CAGGCTAGCGCAGCCATGTGTTCACCAGCAGTTG3'						
5	Reverse 5'CTGCTCGAGCTAATGGTGATGGTGATGGTGACTGCAG						
6	GGCACAGATG3 '						
7	Reverse 5'CTGCTCGAGCTAACTGCAGGGCACAGATG3						
8							
9	The DNA sequences of the above primers are provided						
10	as Sequence ID No's 1 to 5 in the Sequence Listing						
11	Section of this specification.						
12	•						
13	The RT-PCR reaction mix contained 5 $\mu l$ of 10 $ imes$ HF RT-						
14	PCR buffer, 100 ng of forward primer; 100 ng of						
15	reverse primer, 200 $\mu M$ of dNTP, 100 ng of RNA, 1 U						
16	of StrataScript RT (1 unit), and the Taqplus						
	or beracabetipe ki (i wilt), and the Tapplus						
17	Precision DNA polymerase						
17 18							
18	Precision DNA polymerase						
18 19	Precision DNA polymerase  RT-PCR conditions were:						
18 19 20	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle						
18 19 20 21	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle 95°C 1 min 1 cycle						
18 19 20 21 22	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle  95°C 1 min 1 cycle  95°C 30 sec						
18 19 20 21 22 23	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle  95°C 1 min 1 cycle  95°C 30 sec  55°C 30 sec  55°C 30 sec  30 cycles						
18 19 20 21 22 23 24	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle  95°C 1 min 1 cycle  95°C 30 sec  55°C 30 sec  55°C 30 sec  68°C 2 min  1 cycle  30 cycles						
18 19 20 21 22 23 24 25	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle 95°C 1 min 1 cycle 95°C 30 sec 55°C 30 sec 55°C 30 sec 68°C 2 min 68°C 10 min 1 cycle						
18 19 20 21 22 23 24 25 26	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle 95°C 1 min 1 cycle 95°C 30 sec 55°C 30 sec 55°C 30 sec 68°C 2 min 68°C 10 min 1 cycle						
18 19 20 21 22 23 24 25 26 27	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle 95°C 1 min 1 cycle 95°C 30 sec 55°C 30 sec 55°C 30 sec 68°C 2 min 68°C 10 min 1 cycle 4°C ∞  The RT-PCR products were analyzed by means of 1.5%						
18 19 20 21 22 23 24 25 26 27 28	Precision DNA polymerase  RT-PCR conditions were:  42°C 30 min 1 cycle  95°C 1 min 1 cycle  95°C 30 sec  55°C 30 sec  55°C 30 sec  68°C 2 min  68°C 10 min 1 cycle  4°C ∞						

visualized on an UV transiluminator.

32

31

32

1	Amplification of the $\alpha$ and $\beta$ -chains of IL-12
2	starting from the cDNAs
3	
4	The cDNAs coding for the $\beta$ -chain (p40) and $\alpha$ -chain
5	(p35) of interleukin-12 were obtained from ATTC
6	(American Type Tissue Culture Collection, N 40854)
7	and HGMP Resource Centre (Human genome mapping
8	project, Image Clone 1932948, www.hgmp.mrc.ac.uk),
9	respectively. Pwo DNA polymerase from Boehringer
10	Mannhein was the enzyme used for amplification. This
11	enzyme has 3'-5' exonuclease proofreading activity.
12	Amplification was performed for 20 cycles (1 min at
13	95°C, 1 min at 47°C and 1 min at 72°C), using
14	different concentrations of MgSO <sub>4</sub> (2, 3 and 4 mM),
15	200 µM dNTP (Pharmacia), 600 nM of each primer and
16	50 ng of template DNA. A Bio-Rad thermocycler was
17	used for amplification of these products, and the
18	primers used were the same as indicated above.
19	
20	Purification of PCR products
21	
22	PCR products were purified by means of
23	phenol/chloroform extraction. An identical volume of
24	phenol/chloroform/isoamyl alcohol (25:24:1 v/v/v)
25	was added to the samples. Samples were vortexed for
26	1 min and centrifuged at 18,000 rpm for 3 min, in
27	order to separate the different phases.
28	Subsequently, the aqueous phase was collected
29	carefully. We removed the primers with cleaning

columns from QIAGEN. As an alternative to the use of

QIAGEN columns, ethanol precipitation was performed

by adding 3 volumes of ethanol to the samples. 1/10

G-N-N

- volume of sodium acetate (pH=5) was added to the 1
- reactions. Samples were left at -20° C for 1 hour, 2
- and a DNA pellet was obtained by centrifugation at 3
- 4 18,000 rpm for 10 min at 4° C. Pellets were washed
- two times with 1 ml of 70% ethanol to remove salt 5
- and any organic molecules. The pellet was dried at 6
- room temperature and resuspended in 15  $\mu l$  of TE 7
- 8 buffer.

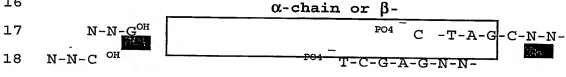
9

#### Restriction digestion of the $\alpha$ and $\beta$ -chains 10

11

- The PCR products were digested with the restriction 12
- enzymes NheI and XhoI which recognise the sequences 13
- $G\downarrow CTAGC$  and  $C\downarrow TCGAG$ , respectively. Both restriction 14
- endonucleases were supplied by Amersham Pharmacia. 15

16



19 -N-N-C-G-A-T OH

20 N-N-G-T-G-C-T OH C-N-N-

21

- One µl of each enzyme (8 and 9 units respectively) 23
- and 2 µl of 10x OPA+ (One-Phor-All Buffer Plus) 24
- buffer were added to 16 µl of purified PCR product, 25
- to make up a final volume of 20  $\mu$ l. The reactions 26
- were incubated at 37°C for 1.5 hours. The digestion 27
- was finalized by heat inactivation of the enzyme 28
- 29 during 20 minutes at 65°C followed by incubation at
- 30 room temperature for 20 min. To concentrate the
- digestion products by precipitation, 1/10 volume of 31

1 sodium acetate (pH=5) and ethanol were added to the

- 2 reactions. Samples were left at -20°C for 1 hour,
- 3 and the pellet was obtained by centrifugation at
- 4 18,000 rpm for 10 min at 4°C. The pellet was washed
- 5 2 times with 1 ml of 70% ethanol. The pellet was
- 6 allowed to dry at room temperature and resuspended
- 7 in 15 µl of TE buffer.

8

- 9 The purified PCR products were subjected to 1.5%
- 10 agarose gel electrophoresis in TBE buffer (45  ${
  m mM}$
- 11 Tris-Borate, 1 mM EDTA) and the bands (700 bp for lpha-
- 12 chain and 900 bp for  $\beta$ -chain) were visualized after
- 13 staining in TBE buffer supplemented with 0.5 µg/ml
- 14 ethidium bromide (30 min) on a UV trans-illuminator.

15

- 16 Restriction digestion of pIND, pIND(SP1) and
- 17 pINDHygro vectors

- 19 The pIND, pIND(SP1) and pINDHygro vectors (ecdysome-
- 20 inducible mammalian expression vectors) were
- 21 supplied by Invitrogen. These vectors each contain
- 22 an ampicillin resistance gene for selection in E.
- 23 coli cells, and either a neomycin (only pIND and
- 24 pIND(SP1)) or an hygromycin resistance gene
- 25 (pINDHygro) for selection in mammalian cells. 2 µg
- 26 of each vector were digested with 8 units of NheI
- 27 and 9 units of XhoI, in 1x OPA buffer in a final
- 28 volume of 20  $\mu$ l. Reactions were incubated at 37°C
- 29 for 1.5 hours and heat-inactivated at 65°C for 20
- 30 min. The vector DNA was precipitated as described
- 31 above.

1 Ligation of the  $\alpha$ -chain into pIND and pINDSP1, and

2 of the β-chain into pINDSP1 and pINDHygro

3

- 4 Ligation of the digested PCR products ( $\alpha$  and  $\beta-$
- 5 chains) into digested vectors was catalyzed by  ${f T_4}$
- 6 DNA ligase enzyme (Promega). Two different ratios of
- 7 vector/insert (1:3 and 1:6) were tested in order to
- 8 optimize the ligation reaction. The reactions were
- 9 performed in a final volume of 20  $\mu$ l, containing 2
- 10  $\mu$ l of 10imes T<sub>4</sub> ligase buffer, 1.5 units of T<sub>4</sub> DNA
- 11 ligase, 3  $\mu$ l of vector (100 ng), and the insert and
- 12 vector DNA. The reactions were incubated overnight
- 13 at 16°C.

14

15 Preparation of competent cells

- 17 E. coli JM109 (endA1, recA1, gyrA96, thi, hsdR17
- 18  $(r_k-, m_k+)$ , relA1, supE44,  $\Delta(lac-proAB)$ , [F',
- 19 traD36, proAB, lacI $^{
  m q}$ Z $\Delta$ M15] cells were made competent
- 20 by means of the  $CaCl_2$  method (REF). A single clone
- 21 was inoculated in 5 ml of LB (Luria-Bertani broth
- 22 containing 10 g/l bactotryptone, 5 g/l bacto-yeast
- 23 extract and 10 g/l NaCl) medium and left overnight
- 24 with vigorously shaking at 37°C in a dedicated
- 25 incubator. An aliquot of this culture (100  $\mu$ l) was
- 26 added to 5 ml of LB (Luria B) medium. This culture
- 27 was further incubated at 37°C until an OD  $(A_{600})$  of
- 28 0.5 was reached (log phase). Cells were placed on
- 29 ice for 5 minutes and then distributed (1 ml) in
- 30 sterilized eppendorf tubes. These tubes were
- 31 centrifuged at 13,000 rpm for 5 minutes,
- 32 supernatants were discarded and pellets were

- 1 resuspended in 1 ml of ice-cold CaCl2. The cells
- 2 were pelleted by centrifugation at 13,000 rpm for 5
- 3 minutes at 4°C, and washed in 1 ml of ice-cold
- 4 CaCl2; the pellet obtained was now resuspended in
- 5 200  $\mu$ l of CaCl<sub>2</sub> and frozen at -70°C.

### 7 Transformation of E. coli cells

8

- 9 Transformation was performed by mixing an aliquot of
- 10 competent cells with the ligation reactions (7.5
- 11  $\mu$ l). This mixture was incubated on ice for 1 hour
- 12 and then subjected to a heat-shock at 42°C for 2
- 13 minutes. 1 ml of LB medium was added, and this
- 14 suspension was left at 37°C for 1 hour with
- 15 vigorously shaking. The transformation reactions
- 16 were mixed with 0.7 % agar supplemented with 50
- 17  $\mu$ g/ml ampicillin and then plated on preheated (37°C)
- 18 LB 1.5 % agar plates containing ampicillin (50
- 19  $\mu$ g/ml). The plates were incubated overnight in an
- 20 incubator at 37°C.

21

# 22 Plamid purification from transformed E. coli cells

- 24 Colonies were inoculated in 5 ml of LB medium
- 25 containing 50 μg/ml of ampicillin and left overnight
- 26 with vigorously shaking at 37°C in an incubator.
- 27 Cells were collected by centrifugation at 6,000 rpm
- 28 for 5 min. Pelleted cells were processed with the
- 29 Qiagen miniprep purification kit. Qiagen plamid
- 30 purification kits are based on an alkaline lysis
- 31 procedure using a buffer composed of SDS, that
- 32 disrupt the cell membranes, and NaOH, known to

- 1 denature genomic DNA. The cell lysate is loaded onto
- 2 an anion exchange resin that captures the DNA.
- 3 Afterwards, RNA, proteins, dye and impurities are
- 4 removed with a medium salt buffer (1 M NaCl). DNA is
- 5 eluted by means of a buffer that contains 1.25  ${\tt M}$
- 6 NaCl. The eluted DNA is concentrated and
- 7 precipitated with isopropanol.

- 9 Sequencing of pIND(SP1)-p35H, pIND-p35H, pIND-40H,
- 10 pINDHygro-p40, pIND(SP1)-p40H and pIND-p40

11

- 12 The sequence of inserts was verified by the
- 13 enzymatic dideoxy-method described by Sanger et al.
- 14 (1977). The 'Ecdysone Forward' and 'BGH Reverse'
- 15 primers were used for forward and reverse
- 16 sequencing, respectively. The ABI PRISM Big DYE
- 17 Terminator Cycle Sequencing Ready Reaction Kit was
- 18 used. A mixture was prepared consisting of 8 µl of
- 19 the Terminator Ready Reaction Mix, 3.2 pmol of each
- 20 primer and 500 ng of DNA, and deionized water was
- 21 added to a volume of 20  $\mu$ l. PCR conditions were 25
- 22 cycles 15 sec at 50°C, 25cycles
- 23 60°C for 4min
- 24 4°C ∞

- 26 Prior to sequencing, PCR products were purified in
- 27 order to remove dNTPs, primers and unincorporated
- 28 dye terminators. Ethanol precipitation was carried
- 29 out by adding 2  $\mu$ l of 3 M sodium acetate pH=4.6, and
- 30 50  $\mu$ l of 95 % ethanol to the PCR products. Samples
- 31 were vortexed and left at room temperature for 15
- 32 minutes. Subsequently, the samples were centrifuged

- at 18,000 rpm (4°C) for 20 minutes. The supernatant 1
- fractions were discarded and the pellet was washed 2
- two times with 270  $\mu l$  of 70 % ethanol. The pellet 3
- 4 dried at room temperature, followed
- resuspension in 5  $\mu$ l deionized formamide and 25 mM 5
- EDTA to which blue dextran was added (50 mg/ml). The 6
- samples were heated at 95°C for 2 minutes before 7
- being loaded on an ABI PRISM 310 Genetic Analyzer. 8

9

#### 10 Cell cultivation and transfection

11

#### 12 Maintenance of cells

13

- The human embryonic kidney cell line (EcR-293), 14
- previously transfected with a pVgRXR construct that 15
- 16 encodes the regulatory ecdysone receptor,
- obtained from Invitrogen. The cells were cultured in 17
- DMEM (LifeTechnologies) supplemented with 10 % of 18
- 19 foetal bovine serum (LifeTechnologies)
- glutamine 2 mM, in addition to 400  $\mu g/ml$  zeocin, 400 20
- $\mu g/ml$  hygromycin or 600  $\mu g/ml$  G418 for selection of 21
- 22 transfected cells (Invitrogen). Cells were
- cultivated in 75 cm<sup>2</sup> flasks until 80% of confluency 23
- was reached. Medium was removed and trypsin-EDTA 24
- solution was added. After 15 minutes at 37 C, medium 25
- was added and cells were collected. The suspensions 26 27
- were centrifuged at 1,000 rpm for 5 min. in order to remove the trypsin. Cells were resuspended in medium 28
- and transferred to new culture flasks. Cells were 29
- generally split 1 over 10 once a week. Cells were 30
- maintained in a  $CO_2$  incubator at 37°C (5%  $CO_2$ ). 31

1	Freezing	<u>of</u>	EcR-293	clones	expressing	IL-12	α	or
---	----------	-----------	---------	--------	------------	-------	---	----

2 β-chains

3

4 Selected clones were cultivated in 175 cm2-flasks until they reached 80 % confluency. The cells were 5 collected by trypsinization, 6 and counted in 7 hemacytometer by means of the trypan blue exclusion assay - REF ). Cells were resuspended at a density 8 of  $3\times10^6$  cells/ml in the freezing medium, which was 9 composed of 90 % medium and 10% DMSO, and these 10 11 suspensions were transferred to cryovials. cryovials (LifeTechnologies) were placed at -20°C 12 for 2 hours, transferred to a -70°C freezer for 16

13

hours and, finally, placed in liquid nitrogen for 14

15 long-term storage.

16

#### 17 Transfection of mammalian cells

18

Plasmid DNA used for transfection of mammalian cells 19

was purified by means of the Endofree kit of QIAGEN. 20

21 The purified plasmid DNA was quantified by

22 spectrophotometry. DNA concentrations were

determined by measuring absorbance at 260 nm, 23

the purity was estimated by the  ${\rm A}_{\rm 260}/{\rm A}_{\rm 280}$  ratio. 24

25

EcR293 cells were plated in 6-well plates  $(2\times10^5)$ 26

the day before the transfection. Transfections of 27

EcR293 cells were performed by means of the FuGENE-6 28

transfection reagent (Boehringer Mannheim). FuGENE-6 29

is a cationic lipid reagent which interacts with 30

negatively charged DNA to form a complex that can 31

cross the cell membrane. We used 1 or 2  $\mu g$  of 32

- 1 plasmid DNA (pIND(SP1)-p35H, pINDHygro-p40 or pIND-
- 2 p40H) to transfect cells. DNA samples were mixed
- 3 with 3  $\mu l$  of FuGENE-6, and diluted in 97  $\mu l$  of
- 4 medium. This solution was directly added to the
- 5 cells.

- 7 Preparation of soluble and insoluble fraction of
- 8 cells

9

- 10 Monolayers of EcR293 cells were washed 3 times with
- 11 large volumes of PBS. Cells were scraped and
- 12 resuspended in PBS, and centrifuged. The pelleted
- 13 cells were resuspended in lysis buffer, and
- 14 incubated on ice for 30 minutes. Lysis buffer was
- 15 composed of PBS, supplemented with 5 mM EDTA, 5 mM
- 16 EGTA, 1xprotease inhibitors (Boehringer Mannheim),
- 17 and 1% Triton X-100. Subsequently, the samples were
- 18 centrifuged at 18,000 rpm for 10 minutes, and the
- 19 soluble fraction recovered. The insoluble fraction
- 20 was washed with PBS supplemented with 1% Triton X-
- 21 100, and centrifuged at 18,000 rpm for 10 minutes.
- 22 Both the soluble and insoluble fractions were now
- 23 ready for analysis by SDS-PAGE and immunoblot.

24

25 Gel electrophoresis (SDS-PAGE)

- 27 Sodium dodecyl sulphate polyacrylamide
- 28 electrophoresis (SDS-PAGE; Laemmli, 1970) was used
- 29 as a standard technique for separating proteins in
- 30 the culture medium, soluble/insoluble cell
- 31 fractions, and immunoprecipitates. Generally,
- 32 protein samples were mixed with 2x SDS-PAGE loading

- 1 solution and loaded into the wells of pre-cast 4-15%
- 2 polyacrylamide gels. Electrophoresis was performed
- 3 at high voltage (200V) using a BioRad Mini-Protean
- 4 III electrophoresis unit and a Pharmacia power
- 5 supply. The electrophoresis buffer used contained 25
- 6 mM Tris, 192 mM glycine, and 0.1 % SDS (pH=8.3).
- 7 Size standards, such as the 'Perfect Protein Western
- 8 Blot Marker' from Novagen, were included in every
- 9 gel.

11 Western blotting, antibodies and detection

12

13 Immunoblot

- 15 Following SDS-PAGE, proteins were transferred from
- 16 the gel to a PVDF membrane by semi-dry
- 17 electroblotting. The polyacrylamide gel and 2 stacks
- 18 of pre-cut Whatman filter papers were equilibrated
- 19 in transfer buffer (48 mM Tris, 39 mM glycine, 0.04
- 20 % SDS, 20, % methanol) for 10 minutes. A PVDF
- 21 membrane was briefly soaked in methanol. The gel and
- 22 the PVDF membrane were placed between two stacks of
- 23 ten layers of filter papers, and the whole was
- 24 transferred to an electro-blotting unit. The
- 25 electrotransfer conditions applied were  $0.8~\mathrm{mA/cm^2}$
- 26 for 1 hour. The apparatus was dismantled, and the
- 27 membrane was incubated overnight at 4°C in blocking
- 28 buffer (2 % casein in TBS consisting of 10 mM Tris-
- 29 HCl, pH=7.4, and 100 mM NaCl). The membrane was
- 30 incubated with a primary antibody. We used the
- 31 following antibodies: (i) mouse  $\alpha$ -p35 antibody G161-
- 32 566, obtained from BD-PharMingen, and used at a

- 1 working concentration amounting to 1/10,000 of the
- 2 original stock; (ii) mouse  $\alpha$ -p40 antibody C8.6, BD-
- 3 PharMingen, used at a 1/5,000 dilution; or (iii) the
- 4 mouse anti-IL-12 antibody 1-2A1 obtained from Abcam,
- 5 1/1,000 diluted. For detection of chaperones we used
- 6 the following antibodies: (i) anti calreticulin, and
- 7 (ii) anti GR894, from Stratogen.

- 9 These primary antibodies were added to TBS-T, i.e.
- 10 TBS supplemented with 0.5% Tween-20 and 0.1 %
- 11 casein. Incubation was done at room temperature for
- 12 2 hrs. Membranes were washed repeatedly with TBS-T
- 13 buffer (without casein), and subsequently incubated
- 14 with a secondary antibody. The secondary antibody
- 15 used was either goat anti-mouse or goat anti-rabbit
- 16 horseradish-peroxidase-conjugated antibody from
- 17 Jackson&ImmunoResearch (used at a 1/20,000
- 18 dilution). Incubation was performed for 1 hour at
- 19 room temperature, after which membranes were washed
- 20 again. The 'Perfect Protein Western Blot Marker' was
- 21 detected by means of an S-protein HRP conjugate
- 22 (Novagen), used at a working concentration of
- 23 1/5,000 of the orginal stock. Detection of poly-
- 24 histidine tagged fusion proteins was carried out
- using the  $INDIA^{TM}$  HisProbe-HRP purchased from Pierce.
- 26 In this case, following overnight blocking, the
- 27 membrane was incubated with INDIA HisProbe (1/5,000
- 28 dilution) in TBS-T buffer with 0.1 % casein.

29

30 Chemiluminiscent detection

Chemiluminiscent detection was carried out with 1 either the 'ECL' or 'ECL+Plus' kit , both purchased 2 from Amersham-Pharmacia. The ECL detection principle 3 4 based on the oxidation of luminol diacylhydracide), while ECL+Plus uses the enzymatic 5 6 generation of an acridinium ester. The produces a more intense light emission of longer 7 duration. According to the manufacturer, the ECL kit 8 can generally detect 1 pg of antigen, while the 9 ECL+Plus kit can detect 20 times less protein. When 10 using the ECL kit, the working solution was prepared 11 by mixing equal parts of the 'Luminol/Enhancer' and 12 'Peroxidase' solutions. When using the ECL+Plus kit, 13 the working solution was prepared by mixing 40 parts 14 of the 'Substrate' solution with 1 part of 'Acridan' 15 The membrane was incubated with these 16 solutions for 5 or 1 minute(s), respectively. Excess 17 solution was removed from the membrane. The membrane 18 was wrapped in cling film, and exposed using Kodak 19 20 MR1 or MR2 films.

21

## 22 Stripping and reprobing of membranes

23

24 Primary and secondary antibodies were removed from the membranes by incubation in stripping buffer (100 25 mM 2-mercaptoethanol, 2 % SDS, and 62.5 mM Tris-HCl; 26 pH=6.7). Incubation was allowed to proceed for 30 27 min. to 1 hour at 50-60°C. The membrane was washed 28 in TBS-T for 1 hour and blocked in 2% casein. At 29 this stage, the membrane was ready for re-incubation 30 31 with a primary antibody.

- Purification of the recombinant  $\alpha$  and  $\beta$  subunits of 1 2 IL-12 3 Ni<sup>2+</sup>-NTA chromatography 4 5 Purification of hexahistidine-tagged  $\alpha-$  and  $\beta-$ chains 6 was performed using nickel-nitrilotriacetic acid 7  $(Ni^{2+}-NTA)$  affinity chromatography.  $Ni^{2+}-NTA$  agarose 8 9 was obtained from QIAGEN. 10 11 Cross-linking of proteins 12 Following induction, cells were washed, scraped and 13 resuspended in PBS supplemented with 100  $\mu g/ml$  of 14 dithiobis(succinimidylpropionate (DSP). DSP is a 15 homobifunctional NHS-ester that reacts with the  $\varepsilon\text{--}$ 16 amines of lysines residues, so as to form a covalent 17 amide bond. Cross-linking reactions were incubated 18 19 at room temperature for 30 minutes, with intermittent vortexing performed every 5 minutes. 20 Reactions were quenched by adding 100 mM of Tris.HCl 21 (pH=8.0). As Tris contains DSP-reactive primary 22 amines, the aim of this 'quenching' reaction is to 23 block any remaining unreacted DSP. Quenching was 24 25 allowed to proceed for 15 minutes. 26 Inhibitor and cytotoxicity assays 27 28 29 Inhibitor assay 30
- To analyse the effect of inhibitors on formation and secretion of IL-12, generally cells were grown in

5μΜ

- 1 12-well plates. When the cells reached a confluency
- of 70 %, inhibitors were added to the culture medium 2
- at the concentrations indicated. After 2 hours of 3
- incubation, cells were induced with ponasterone A. 4
- 5 Sixteen to twenty-four hrs later, medium
- collected to analyse secretion of  $\alpha$  and  $\beta$ -chains, 6
- either alone or in combination. Cells were lysed as 7
- described above, and soluble and insoluble fractions 8
- 9 were prepared. In some experiments, the a- and/or b-
- chains were purified by means of Ni2+-NTA agarose 10
- affinity chromatography. 11

12

13		INHIBITION OF	Concentration
14	A23187	Ionophore	0.1 to 30µM
15	CELEBREX	Cox-2 Inhibitor	i0 to 100μM
16	Thapsigargin	ER Ca-ATPase	5µM

17

18

19 Cytotoxicity test

- The mitochondrial ...MTT test is widely use as a 21 cytotoxicity test. This test is principally based on 22 the propensity of mitochondrial dehydrogenases to 23 cleave the tetrazolium ring of. The viability of 24 25 cells is proportional to the activity of 26 mitochondrial dehydrogenases. Cleavage the tetrazolium ring results in the formation of purple 27 formazan crystals. We used the MTT assay to quantify 28 cytotoxicity of celecoxib on EcR293 cells. The test 29 was performed in 96-well plates in which 105 cells 30
- per well were plated the day before application of 31
- the MTT test. Following addition of celecoxib to the 32

- 1 culture medium, cells were induced by ponasterone A,
- 2 as explained before. After 16 hours of induction,
- 3 the MTT reagent (10 µl of 100 mg/ml stock solution)
- 4 was added to the cells. Two hours later, the medium
- 5 was removed, and the cells were dissolved in DMSO.
- 6 DMSO solubilizes formazan crystals. Absorbance was
- 7 measured at 550 nm using a 96-well plate
- 8 spectrophotometer.

- 10 Description of the Ecdysone-Inducible Mammalian
- 11 Expression System

- 13 As a means to study folding and secretion of dimeric
- 14 forms of interleukin, a series of cell lines that
- 15 produce the recombinant  $\alpha$  and  $\beta$ -chain under
- 16 transcriptional control of a chemically inducible
- 17 promotor were developed. The expression system used
- 18 is based on the ability of the insect hormone
- 19 ecdysone (analog Ponasterone A) to induce
- 20 transcription of IL-12 in mammalian cells from a
- 21 compatible promoter. Since mammalian cells do not
- 22 express the ecdysone receptor, the basal levels of
- 23 transcription of IL-12 were low or non-existent. The
- 24 hormone ecdysone (or its analogs) does not affect
- 25 the physiology of mammalian cells, and hence, can be
- 26 used without inducing any other irrelevant or toxic
- 27 effects. This expression system facilitates
- 28 extremely tight control of the expression of  $\alpha$  and
- 29  $\beta$ -chain genes, which is of interest for both kinetic
- 30 studies and studies in which inhibitors are used as
- 31 a means to monitor the process of folding and
- 32 secretion of IL-12.

Architecture and components of the 2 Ecdysone-3 Inducible Mammalian Expression System

4 The Ecdysone-Inducible Mammalian Expression System 5 6 (EIMES) is based on the use of a heterodimer composed of the ecdysone receptor (VgEcR) and the 7 retinoid X receptor (RxR) (Figure 1A). Both receptors 8 are coded for in the cell line by the plasmid pVgRxR 9 vector that carries the zeocin resistance gene, 10 allowing for selection by means of this antibiotic. 11 12 receptor is under ecdysone transcriptional control of the Rous sarcoma virus promoter  $(P_{RSV})$ 13 while the retinoid receptor is located downstream 14 from the cytomegalovirus promoter ( $P_{\text{CMV}}$ ). Both are 15 16 constitutive promoters facilitating continuous production of high levels of the heterodimer. The 17 ecdysone receptor contains the VP16 transactivation 18 domain which increases the level of induction. 19 the presence of ponasterone A (ecdysone analog) the 20 ecdysone and retinoid X receptors will bind to each 21 22 other, and the heterodimerized receptor 23 subsequently bind to the ecdysone/glucocorticoid 24 response element (E/GRE) sequence present in the 25 promoter of pIND vectors to be used as vehicle for 26 expression of IL-12 chains (Figure 1B). 27 receptors have a DNA binding domain (DBD) recognises half of the response element (E/GRE). The 28 29 DBD of the ecdysone receptor recognises 5'AGTGCA3' and the DBD of the retinoid receptor recognises the 30 31 sequence 5*'* AGAACA3' (Yao et al., 1993). response element is upstream from the promoter that 32

- 1 activates gene expression  $(P_{\Delta HSP})$  in pIND. Thus the
- 2 binding of the receptor heterodimer to these
- 3 response elements will induce the transcription of
- 4 the gene of interest (Figure 1B). The cell line used
- 5 is EcR293, a derivative of the HEK293 cell line that
- 6 is transfected with the pVgRXR vector and cultivated
- 7 in the presence of zeocin.

9 pIND expression vectors for production of IL-12

10

- 11 Three different pIND vectors (pIND, pINDSP1 and
- 12 pINDHygro) are available all of which can be used in
- 13 this expression system to produce recombinant
- 14 proteins (Figure 2). All of these contain an
- 15 ampicillin resistance gene to enable selection and
- 16 propagation of clones in E. Coli cells. The multiple
- 17 cloning site is located downstream from a minimal
- 18 heat shock promoter ( $P_{\square HSP}$ ). pIND and pINDSP1 differ
- 19 from pINDHygro in that the first two vectors contain
- 20 the neomycin resistance gene while pINDHygro
- 21 contains the hygromycin resistance gene. These
- 22 different antibiotic resistance genes allow for dual
- 23 selection of transfected cells in the presence of
- 24 both antibiotics. This is important in view of the
- 25 requirement of producing cell lines that express
- 26 both subunits of dimeric interleukins, with each
- 27 subunit provided by a different vector.

- 29 The pINDSP1 vector contains three SP1 binding sites
- 30 inserted between the response elements and the
- 31 promoter, which theoretically increases the

- 1 expression levels five times in comparison with pIND
- 2 (Kadonaga et al., 1987).

4 Rational for use of histidine tags

- 6 The use of the histidine tag as a means for
- 7 purification of recombinant proteins is a well-
- 8 documented method proven to be highly efficient.
- 9 The major advantages of this system are:
- 10 Purification can be achieved from a mix containing
- 11 less than 1 % of total protein in one-step.
- 12 Purification can be completed under native or
- 13 denaturing conditions since the binding of the
- 14 histidines to the Ni-NTA agarose is not dependent on
- 15 the conformation. The His tag is a small tag and it
- 16 does not interfere with the structure or function of
- 17 the protein to be expressed so removal of the tag is
- 18 not necessary. The His tag can be used as the target
- 19 to be recognized by an antibody anti-His tag. The
- 20 histidine tag can be engineered so as to be
- 21 expressed in the target protein in either N-
- 22 (preceded by ATG initiation codon) or C-terminal
- 23 (followed by TAA, TGA or TAG stop codon) position.
- 24 This is accomplished through the use of specific
- 25 primers which are designed so as to contain the
- 26 coding sequence for 6 histidines fused to the
- 27 sequence of our target protein. By means of metal
- 28 ionic affinity chromatography (matrix used Ni2+-
- 29 nitrilotriacetic acid coupled to agarose,
- 30 abbreviated as Ni-NTA) His-tagged recombinant
- 31 proteins can be captured and purified in a highly
- 32 selective and specific manner. This strategy was

- 1 applied to the purification of the IL- lpha and eta-
- 2 chains from both cell lysates (in order to capture
- 3 protein in the process of folding in the endoplasmic
- 4 reticulum and to co-capture proteins associated with
- 5 the folding chains such as chaperones) and medium
- 6 (so as to capture fully folded and matured secreted
- 7 protein).

9 Amplification of  $\alpha$  and  $\beta$  chains of IL-12

10

11 Design of primers

12

- 13 The composition of the nucleotide sequence preceding
- 14 the ATG translation initiation codon is known to
- 15 affect translation initiation. Therefore primers
- 16 optimized for translation were designed (consensus
- 17 sequence: GCCRCC ATG). To clone both subunits
- 18 directionally into the multiple cloning sites of
- 19 pIND plasmids, an NheI restriction site was
- 20 introduced in the forward primers and an XhoI
- 21 restriction site in the reverse primers (Figure 3).
- 22 The  $\alpha$  and  $\beta\text{--chain}$  sequences of IL-12 (Sequence ID
- 23 No.s 6 and 7) (Genbank accession numbers: M65291 and
- 24 M65290) were checked to assure that none of these
- 25 contain these restriction sites.

- 27 The IL-12  $\alpha$ -chain sequence contains two initiation
- 28 codons (ATG), which occur in the same reading frame
- 29 and are 99 nucleotides apart. It has been
- 30 demonstrated that  $\alpha$ -chains translated from either
- 31 the first or second start codon are functional.
- 32 Thus, the initiation codon used may affect the



length of the signal peptide, but does not affect 1 primary structure and folding of the mature chain. 2 3

- This is understandable since folding occurs in the
- ER after the signal peptide has been removed. The 4
- forward primer was designed to contain the second 5
- start codon of the functional  $\alpha$ -chain. The reverse 6
- primer contained the stop codon 7 (TAA) and the
- sequence for six histidines engineered between the 8
- carboxy-terminus and the stop codon. Similarly, the 9
- $\beta$ -chain primers contained ATG and TAG stop codons. 10
- For the  $\boldsymbol{\beta}$  chain, however, two reverse primers were 11
- designed, i.e. one containing the sequence coding 12 .
- for the six histidines and the other without the 13
- 14 histidine tag (Figure 3).

15

- Amplification of the  $\alpha$  and  $\beta$  chains of IL-12 by RT-16
- PCR from U937-extracted mRNA 17

- In order to obtain mRNA of the IL-12  $\alpha$  and  $\beta$  chains, 19
- a monocytic cell line (U937) was induced with LPS 20
- for 16 hours, a treatment which is known to result 21
- in the production of IL-12 in this cell line. The 22 23
- RNA was extracted, and mRNA was retrotranscribed 24
- into cDNA by RT-PCR using the primers described in 25 preceding
- paragraph and the high-fidelity 26 thermostable Pwo DNA polymerase. Since the
- concentration of MgSO4 is known to influence the 27
- specificity of primer annealing three 28 different ·
- 29 concentrations of MgSO<sub>4</sub> were used in the
- reaction. Subsequently, the amplification products 30
- 31 were analysed by means of 1.5% agarose
- electrophoresis. Though a band was visible that 32

- 1 corresponded to the expected length of the amplified
- 2  $\beta$  chain (900 bp; Figure 4), no amplification product
- 3 was obtained for the  $\alpha$  chain (not shown).

- 5 Amplification of the  $\alpha$  and  $\beta$  chains of IL-12 by PCR
- 6 from cDNA

7

- 8 The  $\alpha$  and  $\beta$ -chains were amplified using as template
- 9 the full-length cDNAs obtained from the ATCC and the
- 10 HGMP Resource Centre, respectively. Again, we
- 11 decided to use Pwo DNA polymerase for amplification
- 12 rather than Taq polymerase, since the former
- 13 displays 3' → 5' exonuclease proof-reading
- 14 activity which is known to reduce the accumulation
- 15 of errors in the final PCR product. The reactions
- 16 were carried out as explained in section 2.1.3. The
- 17 PCR products obtained by amplification of the cDNAs
- 18 of the  $\alpha$  and  $\beta$ -chains were analyzed by means of 1.5%
- 19 agarose gel electrophoresis. Figure 5 illustrates
- 20 the amplification of the  $\alpha$ -chain: a PCR product
- 21 corresponding to 700 bp was specifically amplified
- 22 in the presence of 2-3 mM MgSO $_4$ . Figure 6 shows the
- 23 900-bp PCR product obtained following amplification
- 24 of the cDNA of the  $\beta$ -chain.

25

26 Construction of pIND-derived expression vectors

27

28 <u>Introduction</u>

- 30 The PCR products were purified and digested with
- 31 NheI and XhoI, and subsequently cloned into
- 32 NheI/XhoI-cut vectors. 5 different constructs were

- 1 created, i.e. pIND-p35H, pIND(SP1)-p35H, pINDHygro-
- 2 p40, pIND(SP1)-p40H and pIND-p40. The expression
- 3 cassettes for the  $\alpha$  and  $\beta$  chains of IL-12 contained
- 4 within these vectors are specified in Figure 7. As
- 5 explained above, pIND(SP1) and pINDHygro confer
- 6 resistance to different antibiotics, i.e. neomycin
- 7 and hygromycin respectively, when expressed in
- 8 mammalian cells. Thus, expression vectors were
- 9 constructed that would facilitate selection of the
- 10 following stable cell lines:

- 12 1. EcR293 cells expressing the carboxyterminal-
- 13 His-tagged  $\alpha$ -chain selected by the antibiotic
- 14 neomycin (transfected with either pIND-p35H or
- pIND(SP1)-p35H, anticipated to differ only in
- the level of expression);
- 2. EcR293 cells expressing the  $\beta$ -chain selected
- with neomycin (pIND-p40 or pIND(SP1)-p40H,
- 19 differing in level of expression but also in
- 20 the presence or absence of a carboxyterminal
- 21 His-tag);
- 22 3. EcR293 cells expressing the  $\beta$ -chain selected
- with hygromycin (pINDHygro-p40)
- 24 4. EcR293 cells expressing the  $\alpha/\beta$  heterodimer
- 25 selected with both neomycin and hygromycin
- 26 (pINDHygro-p40 and either pIND-p35H or
- 27 pIND(SP1)-p35H).

28

29 Selection and sequencing of clones

- 31 Competent E. coli JM109 cells were transformed with
- 32 these different constructs. Following

transformation, the cells were plated on Petri 1 2 dishes containing LB-agar supplemented with 3 pIND vectors confer resistance ampicillin. ampicillin to E. coli cells that have successfully 4 integrated the plasmid. However, still the presence 5 or absence of an insert in the vector has to be 6 verified. In order to confirm the presence of the 7 insert three complementary methods were adopted. 8 First, colony PCR was performed facilitating the 9 identification of positive clones by means of direct 10 amplification of the insert using  $\alpha$  and  $\beta\text{--chain--}$ 11 specific primers. Second, the presence of the insert 12 13 NheI/XhoI restriction digestion of plasmid minipreps and electrophoresis. Third, forward and 14 reverse sequencing was performed to validate the 15 presence of the insert and the absence of 16 errors. The results of the colony PCR procedure are 17 illustrated in Figures 8 and 9, which show that not 18 every ampicillin-resistant colony 19 appeared contain the insert. 20

21

The positive colonies that were identified in Figure 22 8 and 9 were propagated in LB medium supplemented 23 with ampicillin, and minipreps and glycerol stocks 24 25 To confirm the presence of the were prepared. insert in the plasmid minipreps were digested with 26 NheI and XhoI restriction enzymes and these products 27 were subjected to 1.5% agarose gel electrophoresis 28 29 (Figure 10).

30

31 The third method utilised to verify that the 32 plasmids extracted from ampicillin-resistant clones



1	contained	the	correc	t ins	erts	corresp	onding	to
2	either	α	and	β-chai	ns,	consi	sted	of
3	dideoxynuc	leotic	de DNA	sequ	encin	g. For	ward	and
4	reverse se	quenc:	ing was	perfor	cmed u	sing th	e multi	.ple
5	cloning si	te pr	imers,	i.e. e	cdysor	ne forwa	ard pri	.mer
6	and BGH re	verse	primer.	This	showe	d that	error-f	ree
7	inserts we	ere pr	resent	in the	righ	t orien	ntation	in
8	each of the	e vect	ors.					
^								

9

10 Development of stably transfected EcR293 cell lines

11

12 Extraction of endotoxin-free plasmid DNA to be used

13 for transfection of EcR293 cells

14

15 plasmids were purified using the The Endofree purification kit from QIAGEN. This kit facilitates 16 large-scale extraction of plasmid DNA from 100ml of 17 18 bacterial cultures while efficiently removing 19 endotoxins. Endotoxins are toxic for mammalian cells, and their presence in DNA preparations may 20 decrease transfection efficiency . The DNA of the 21 purified samples was quantified by spectrophotometry 22  $(A_{260})\,.$  The concentrations obtained ranged between 23 0.4 and 2  $\mu g/\mu l$  (Table 1). The purity of DNA samples 24 was calculated by absorption measurements at 260 and 25 26 A ratio  $A_{260/280}$  amounting to 1.8 to 2 indicative for a very high purity. As can be seen in 27 Table 1, both the amounts and purities of 28 plasmid DNA obtained using the Endofree kit were 29 30 highly satisfactory.

- 1 Table 1. Concentration, total amount and purity of
- 2 plasmid DNA extracted from bacterial cultures with
- 3 the Endofree kit

Plasmid	A <sub>260</sub>	Conc.	Total	Ratio
			Amt.	(Purity)
pIND	0.051	0.577µg	115.4µg	1.825
(SP1)		/µl		
-р35Н				
pIND	0.070	2.059µg	411.5µg	1.876
Hygro		/µl		
-p40				
pIND	0.097	0.998µg	199.6µg	1.809
-35H		/µl		
pIND	0.047	0.478µg	95.6µg	2.082
-p40		/µl		
pIND	0.098	1.07µg	214µg	1.89
(SP1)		/µl		
-p40H				

5

## Transfection and selection of EcR293 cells

6 7

17

time.

EcR293 cells were transfected with these vectors, 8 either alone or in combinations. Following 1 day of 9 recovery after transfection, cells were trypsinized, 10 diluted and seeded into 96-well plates. 11 appropriate antibiotics were added to the culture 12 medium to initiate the selection\_process. 13 14 summarized in Table 2, three different cell 15 concentrations and two different antibiotic concentrations were used to perform selection over 16

Vectors and vector combinations used to transfect
EcR293 cells:

4

1

- 5 1-pIND-p35H 3pIND-p40 6-pIND-p35H/pINDHygro-40 6 2-pIND- 4pINDHygro 7-pIND(SP1)-35H/pINDHygro
- 7 (SP1)-p35H -p40 -p40
- 8 5pIND(SP1)
- 9 -p40H

10

- 11 Table 2. Cell and antibiotic concentrations for
- 12 selection of transfected EcR293 cells

Conc. Conc. Conc. Neomycin Hygromycin Zeocin Dilution 1/10 (105 cells/well) 300 µg/ml 300 µg/ml 400 μg/ml 600 µg/ml 600 μg/ml 400 μg/ml 10<sup>6</sup> transfected cells 300 µg/ml 300 µg/ml Dilution 1/100 (104 cells/well) 400 μg/ml 600 µg/ml 600 µg/ml 400 μg/ml 300 µg/ml 300 µg/ml 400 μg/ml Dilution 1/1000 (103 cells/well) 600 μg/ml 600 µg/ml 400 µg/ml

13 For the construct made with the pINDHygro vector

- 14 (pINDHygro-p40), selection was performed in the
- 15 presence of either 300 or 600 μg/ml hygromycin.
- 16 These concentrations were chosen on the basis of the
- 17 concentrations of hygromycin recommended by the
- 18 manufacturer of the pIND series of vectors for
- 19 selection of transfected EcR293 cells (between 200
- 20 and 600µg/ml). Similarly, cells transfected with
- 21 pIND- and pINDSP1-derived vectors were cultivated in
- 22 the presence of either 300 or 600 μg/ml neomycin, as
- 23 recommended. Hygromycin concentration of 200 µg/ml

- 1 was used in all further transfection experiments
- 2 with pINDHygro-p40. After 6 weeks we were able to
- 3 detect about 40 different clones in total, generated
- 4 by transfection with the different constructs and
- 5 selection with the appropriate antibiotics.

- 7 Immunodetection of expression of  $\alpha$  and  $\beta$  chains
- 8 following induction with ponasterone A

9

- 10 As a test in order to evaluate whether these clones
- 11 were able to produce the corresponding recombinant
- 12 proteins, we selected three clones, i.e. 1 single
- 13 clone for pIND-p35H (clone 1A9), 1 for pIND(SP1)-
- 14 p35H (clone 2G10) and 1 for pIND-p40 (clone 3D9).
- 15 These clones were trypsinized and plated into the
- 16 wells of 6-well plates. The cells were induced with
- 17 Ponasterone A (5  $\mu M$ ) for 48 hours. Subsequently, the
- 18 cell culture medium was collected, and the cells
- 19 were lysed. This was done to evaluate the presence
- 20 of the recombinant protein in both secreted and
- 21 intracellular fractions. Culture medium and soluble
- 22 cytoplasmic fractions were subjected to 4-15%
- 23 reducing SDS-PAGE (Figure 11). The proteins were
- 24 transferred by electroblot to a PVDF membrane.
- 25 Immunodetection was performed with anti-IL-12  $\alpha$  or
- 26  $\beta$ -chain antibodies. Immunoreactive bands were
- 27 visualized using a chemoluminiscence-based kit and
- 28 autoradiography films, Kodak BioMax MR films. (ECL
- 29 kit; see sections 2.7).

- 31 This first analysis indicated that p40 is more
- 32 efficiently secreted than p35, as the ratio of

- 1 secreted/intracellular is obviously higher for the
- 2 former. Finally, a band corresponding to the Mr of
- 3 serum albumin was visible in all immunoblots of
- 4 medium fractions (indicated with arrow in Figure 11
- 5 A and B). A similar immunoreactive band was found in
- 6 the medium of uninduced or untransfected cells,
- 7 indicating that this band is unrelated to any of the
- 8 IL-12 chains but is likely visualized following a-
- 9 specific interaction with either the primary or
- 10 secondary antibodies used in these experiments (not
- 11 shown).

- 13 Differences in expression levels in stably
- 14 <u>transfected cell lines</u>

15

- 16 Having demonstrated the inducible expression of
- 17 immunoreactive proteins corresponding to either the
- 18  $\alpha$  or the  $\beta$  chain of IL-12 in some of the EcR293 cell
- 19 clones produced, the expression levels in all of the
- 20 clones were evaluated by means of a similar
- 21 procedure. For this purpose cells, precedingly
- 22 seeded in 96 well plates  $(5x10^4 \text{ cells})$  were induced
- 23 with ponasterone A for 24 hours. Induced and
- 24 uninduced cells were lysed in 6 µl of lysis buffer,
- 25 and the lysates were subjected to 4-15% reducing
- 26 SDS-PAGE and immunoblot (Figure 12 and 13).

- .28 Surprisingly, an anti- $\alpha$ -chain reactive band was
- 29 observed in the lysates of both un-induced and
- 30 induced EcR293 cells that exhibited a slightly lower
- 31 Mr than the inducible, recombinant  $\alpha$  -chain. This
- 32 band was also consistently observed in immunoblots

- 1 of un-transfected EcR293 cells (not shown). Thus,
- 2 this protein is likely to correspond to a natural,
- 3 constitutively produced form of either p35 or a p35-
- 4 related protein in these cells. Its Mr is smaller
- 5 than that of the recombinant form, which is likely
- 6 due to the absence of the hexahistidine-tag in the
- 7 natural form. Nevertheless, the smaller form is
- 8 unlikely to correspond to a proteolytically
- 9 generated truncated form of the recombinant his-
- 10 tagged  $\alpha$ -chain as it is equally present in un-
- 11 induced or un-transfected cells.

- 13 Most of the cell lines were freezed and kept in
- 14 liquid nitrogen. Cell line 2B9 (Figure 12, lane 1-
- 15 2), which appeared to be the cell line with the
- 16 highest expression level of the lpha-chain was
- 17 maintained in cultivation for further experiments.
- 18 This cell line was re-named HACHIE.1. Similarly,
- 19 cell line 3H10 which expresses high levels of the  $\beta-$
- 20 chain (Figure 13B, lane 1-2) was maintained in
- 21 culture. This cell line was re-named HIBERNIA.1.

22

- 23 Transient transfection of HIBERNIA.1 cells to
- 24 produce heterodimeric IL-12

- 26 As described above, HIBERNIA.1 is a cell line that
- 27 produces high levels of carboxyterminally
- 28 hexahistidine-tagged  $\beta$ -chain upon induction with
- 29 ponasterone A, and was obtained by transfection of
- 30 EcR293 cells with pIND(SP1)-p40H followed by
- 31 selection with neomycin. The transient transfection
- 32 was carried out in 6-well plates using 1 or 2  $\mu g$  of

- 1 endotoxin-free pIND(SP1)-p35H plasmid DNA. Cell
- 2 culture medium was collected at 30 and 48 hours
- 3 following induction. The samples were run in a non-
- 4 reducing gel so as to facilitate detection of the
- 5 disulfide-bonded heterodimer. Following
- 6 electrophoresis, semi-dry blotting was performed,
- 7 and the membrane was successively probed with an
- 8 anti- $\beta$ -chain (Figure 14) and an anti- $\alpha$ -chain
- 9 antibody (Figure 14).

- 11 Figure 14 shows that in the culture medium of both
- 12 the transiently transfected (lanes 1 to 4) and not-
- 13 transfected (lane 5) HIBERNIA.1 cells 2 immuno-
- 14 reactive bands are detected with the anti- $\beta$ -chain
- 15 antibody, with Mr's of about 40 and 80 kD
- 16 respectively. In lanes 1 to 4, the 80-kD band could
- 17 represent the  $\beta$  chain homodimer (2x40 kD) as well
- 18 as the  $\alpha/\beta$  chain heterodimer (35+40 kD), as both
- 19 would migrate as bands with similar Mr in this low-
- 20 resolution SDS-PA gel. In not-transfected HIBERNIA.1
- 21 cells (lane 5 of Figure 14) the 80 kD band must
- 22 necessarily represent the  $\beta$  chain homodimer. Figure
- 23 14 shows that a 80-kD protein band which is reactive
- 24 with the anti- $\alpha$ -chain antibody is present only in
- 25 HIBERNIA.1 cells transfected with pIND(SP1)-p35H
- 26 (lanes 1 to 4) but not in un-transfected HIBERNIA.1
- 27 cells (lane 5). Analysis of recombinant cell lines
- 28 secreting the  $\alpha$  chain by means of non-reducing SDS-
- 29 PAGE showed that the lpha chain is present only as a
- 30 monomer form when expressed in the absence of the  $\boldsymbol{\beta}$
- 31 chain (data not shown). In view of these findings,
- 32 it can be safely concluded that HIBERNIA.1 cells

transiently transfected with pIND(SP1)-p35H secrete 1 the  $\alpha/\beta$  disulfide-bonded IL-12 heterodimer upon 2 induction with ponasterone A. In fact, in these 3 cells the total amount of  $\boldsymbol{\alpha}$  chain secreted ends up 4 as subunit of the heterodimer form, as anti- $\alpha$ -chain 5 reactivity is only visible as an 80-kD band and not 6 as a 35-kD band. However, it is likely that a 7 8 certain fraction of the β chain produced transiently transfected HIBERNIA.1 cells will still 9 10 present as homodimer. This possibility difficult to exclude in view of the fact that the 11 non-transfected HIBERNIA.1 cells 12 produce the

14

homodimer.

13

Transfection of HIBERNIA.1 cells with with 1 µg 15 16 pIND(SP1)-p35H resulted in a production/secretion of the heterodimer compared to 17 transfection with 2  $\mu g$ . This might be related to the 18 fact that due to the 1:1 stoichiometry of  $\alpha$  and  $\beta$ 19 chain interaction in the heterodimer, a level of  $\alpha$ -20 chain production which is higher than that of the  $\beta$ 21 22 may be counterproductive for efficient formation of the heterodimer. 23

24

To verify the composition of the 80-kD band secreted by transiently transfected HIBERNIA.1 cells, we run the medium collected at 48 hrs after induction from HIBERNIA.1 cells transfected with 1  $\mu$ g of pIND(SP1)- p35H (\* in Figure 14), again, this time in a reducing gel. Gels were blotted, and detection was carried out with either the anti- $\alpha$ -chain antibody,

- 1 the anti- $\beta$ -chain antibody or with both antibodies at
- 2 the same time.

- 4 The anti- $\alpha$ -chain antibody detected a band
- 5 corresponding to 35 kD, while the anti- $\beta$ -chain
- 6 antibody detected a band of approximately 40 kD
- 7 (Figure 15). Thus, the Mr's of the  $\alpha$  and  $\beta$  chains
- 8 produced in transiently transfected HIBERNIA.1 cells
- 9 coincide with those theoretically predicted. The  $\alpha$
- 10 chain appeared as a more diffuse band than the eta
- 11 chain. This is most likely due to more extensive
- 12 heterogeneity in N-glycosylation of the former, as
- 13 tunicamycin treatment produced a much sharper  $\alpha$ -
- 14 chain band (demonstrated below).

15

- 16 This data shows that a genuinely processed  $\alpha$ -chain
- 17 form is produced in transiently transfected
- 18 HIBERNIA.1 cells that interacts with the  $\beta$ -chain to
- 19 form a disulfide-linked secreted IL-12 heterodimer.
- 20 Obviously, these experiments show that attachment of
- 21 hexahistidine-tags to the carboxytermini of both the
- 22  $\alpha$  and  $\beta$ -chains does not interfere with correct
- 23 folding, assembly and secretion of the heterodimer.

24

- 25 Capture of  $\alpha/\beta$  and  $\beta/\beta$ IL-12-H6-chaperone complexes
- 26 on Ni<sup>2+</sup>-NTA

- 28 Following induction with Ponasterone A, cells were
- 29 lysed.  $\alpha/\beta$  and  $\beta/\beta$  -H<sub>6</sub>-chaperone complexes were
- 30 captured on  $Ni^{2+}$ -NTA agarose. The gel was washed 5
- 31 times with buffer A (100mM NaH2PO4, 10mM TrisHCl, 8M
- 32 urea, pH 6.3), and elution was carried out with

- 1 buffer B (same as Buffer A, but pH 4.3). Complexes
- 2 were boiled in SDS loading solution + DTT. Proteins
- 3 were separated by 4-15% SDS-PAGE and transferred to
- 4 PVDF membranes. Detection was carried out using
- 5 anti-p35 antibody G161-566.14 (Pharmingen).
- 6 Membranes were stripped and re-probed successively
- 7 with anti-chaperone antibodies ( $\alpha$  -CRT,  $\alpha$  -Grp78,  $\alpha$
- 8 -Grp94 &  $\alpha$  -CNX; StressGen).

## 10 Experimental findings

- 12 IL-12 is a secretory protein. Secretory proteins are
- 13 defined as proteins that are released by cells into
- 14 the extracellular milieu, and that exert their
- 15 biological activity by binding onto a specific
- 16 membrane receptor of target cells. 'Folding' (i.e.
- 17 generation of a correct three-dimensional structure)
- 18 of secretory proteins, such as IL-12, typically
- 19 occurs in a membrane-surrounded cell organelle,
- 20 named the endoplasmic reticulum (ER). The ER is
- 21 specifically enriched in chaperones, thioredoxin-
- 22 type isomerases and proteins involved in
- 23 glycosylation pathways. An important role of these
- 24 factors is to assist in ensuring correct folding of
- 25 secretory proteins during their transit in the ER
- 26 prior to their secretion into the extracellular
- 27 milieu. Improperly folded secretory proteins are
- 28 generally retained in the ER and subsequently
- 29 degraded by proteases and components of the
- 30 cytosolic proteasome. It was hypothesised that the
- 31 use of selected pharmacological agents that
- 32 interfere with the proper functioning of 'folding'-

- 1 assisting factors in the ER could be used to inhibit
- 2 proper folding, and, hence, secretion of IL-12.
- 3 As a first step, different tightly controlled
- 4 ecdysone-inducible recombinant cell lines expressing
- 5 functional C-terminally hexahistidine-tagged IL-12
- 6  $\alpha/\beta$  (heterodimer) and IL-12  $\beta/\beta$  (homodimer) chains
- 7 were developed. The use of such recombinant cell
- 8 lines alleviates some of the problems related to the
- 9 use of natural producer cells of IL-12 (e.g.
- 10 restricted availability, lack of reproducibility
- 11 etc). These recombinant cell lines were used as a
- 12 means to study the processes that determine
- 13 regulation of folding, assembly and secretion of IL-
- 14 12 homo- and heterodimers. The following inhibitors
- 15 were used: (i) thapsigargin (an ER Ca2+-ATPase
- 16 inhibitor), and (ii) the ionophore A23187 and (iii)
- 17 celecoxib (a putative ER Ca2+ perturbating reagent),
- 18 each over a wide range of concentrations.

- 20 Following a 16-hr treatment of cells with these
- 21 inhibitors, culture medium was collected and the
- 22 presence of secreted IL-12 forms was detected by
- 23 means of non-reducing SDS-PAGE and western
- 24 immunoblot. It was found that neither the  $\alpha/\beta$  nor
- 25 the  $\beta/\beta$  dimer forms of IL-12 were present in the
- 26 culture medium of cells treated with thapsigargin
- 27 when this was added over a concentration range of
- 28 0.1  $\mu$ M to 15  $\mu$ M. The amount of extracellularly
- 29 secreted IL-12 dimer forms produced by thapsigargin-
- 30 treated cells was <5% of that produced by untreated
- 31 cells (maximal suppression was observed for all
- 32 concentrations of thapsigargin greater than or equal

- 1 to 0.1  $\mu M$ ). Similarly, the calcium ionophore A23187
- 2 suppressed formaton of secreted IL-12 dimer forms
- 3 when it was used over a concentration range of 0.1
- 4  $\mu M$  to 30  $\mu M$ , with maximal suppression (>95% compared
- 5 to untreated cells) from 1 μM. Toxicity conferred by
- 6 these inhibitors over the test period of 16 hr as
- 7 measured with the MTT test was observed for
- 8 concentrations of thapsigargin >5-10 µM and for
- 9 concentrations of A23187 >10  $\mu M$ . Thus, the maximal
- 10 suppression of secreted IL-12 dimer production is
- 11 achieved at an inhibitor concentration at which
- 12 toxic effects are totally absent, showing that both
- 13 IL-12-suppressive and cell-toxic effects conferred
- 14 by these inhibitors are independent. Secretion of
- 15 IL-12  $\alpha$  and  $\beta$  monomer forms was suppressed by
- 16 neither thapsigargin nor A23187.

- 18 Both thapsigargin and A23187 are likely to exert
- 19 these effects by decreasing the concentration of Ca2+
- 20 in the ER. It is likely that the resulting
- 21 suboptimal concentration of Ca2+ in the ER blocks the
- 22 activity of Ca2+-dependent chaperones and folding-
- 23 assisting proteins involved in the dimer formation
- 24 of IL-12. It was investigated whether CELECOXIB can
- 25 be used to suppress production of secreted IL-12
- 26 dimer forms.

- 28 Celecoxib was dissolved in DMSO and added to
- 29 recombinant HEK293 cells over a concentration range
- 30 from 10  $\mu\text{M}$  to 100  $\mu\text{M}$ . As a control DMSO-only treated
- 31 cells were used. Celecoxib concentrations were
- 32 chosen on the basis of available literature data,

- 1 and coincide with optimal activity of the compound
- 2 in various cell-based systems. Two hours later cells
- 3 were induced with Ponasterone A to produce IL-12  $\alpha/\beta$
- 4 or  $\beta/\beta$  dimer forms. After 16 hrs of additional
- 5 incubation, culture medium was collected and
- 6 assessed for the presence of IL-12 dimer forms by
- 7 means of non-reducing SDS-PAGE and immunoblot. This
- 8 showed that Celecoxib suppressed production of
- 9 secreted IL-12  $\beta/\beta$  homodimers by >95% when used at a
- 10 concentration equal to or larger than 30  $\mu\text{M}$ ; and of
- 11 secreted IL-12  $\alpha/\beta$  heterodimers by >95% when used
- 12 at a concentration equal to or larger than 10  $\mu M$ .
- 13 Secretion of IL-12  $\alpha$  and  $\beta$  monomer forms was not
- 14 suppressed by Celecoxib. Toxicity as measured with
- 15 the MTT assay was visible when cells were treated
- 16 for 16 hrs with a concentration of Celecoxib equal
- 17 to or larger than 100  $\mu M$ .

- 19 The present data demonstrates that Celecoxib
- 20 efficiently suppresses secretion of IL-12  $\alpha/\beta$  and
- 21  $\beta/\beta$  dimer forms by a post-transcriptional and post-
- 22 translational mechanism that involves Ca2+-dependent
- 23 intracellular retention of IL-12 dimers. Maximal IL-
- 24 12-suppressive effects are observed at a
- 25 physiological Celecoxib concentration in the absence
- 26 of any obvious toxic effects.

- 28 For oral administration, the medicament according to
- 29 the invention may be in the form of, for example, a
- 30 tablet, capsule suspension or liquid. The medicament
- 31 is preferably made in the form of a dosage unit
- 32 containing a particular amount of the active

- 1 ingredient. Examples of such dosage units are
- 2 capsules, tablets, powders, granules or a
- 3 suspension, with conventional additives such as
- 4 lactose, mannitol, corn starch or potatoes starch;
- 5 with binders such as crystalline cellulose,
- 6 cellulose derivatives, acacia, corn starch or
- 7 gelatins; with disintegrators such as corn starch,
- 8 potaote starch or sodium carboxymethyl-cellulose;
- 9 and with lubricants such as talc or magnesium
- 10 stearate. The active ingredient may also be
- 11 administered by injection as a composition wherein,
- 12 for example, saline, dextrose or water may be used
- 13 as a suitable carrier.

- 15 For intravenous, intramuscular, subcutaneous, or
- 16 intraperitioneal administration, the compound may be
- 17 combined with a sterile aqueous solution which is
- 18 preferably isotonic with the blood of the recipient.
- 19 Such formulations may be prepared by dissolving
- 20 solid active ingredient in water containing
- 21 physiologically compatible substances such as sodium
- 22 chloride, glycine, and the like, and having a
- 23 buffered pH compatible with physiological conditions
- 24 to produce an aqueous solution, and rendering said
- 25 solution sterile. The formulations may be present in
- 26 unit or multi-dose containers such as seated
- 27 ampoules or vials.

- 29 If the inflammatory disease is localized in the G.I.
- 30 tract, the compound may be formulated with acid-
- 31 stable, base-liable coatings known in the art which
- 32 began to dissolve in the high pH intestine.

- 1 Formulations to enhance local pharmacologic effects
- 2 and reduce systemic uptake are preferred.

- 4 Formulations suitable for administration
- 5 conveniently comprise a sterile aqueous preparation
- 6 of the active compound which is preferably made
- 7 isotonic. Preparations for injections may also be
- 8 formulated by suspending or emulsifying the
- 9 compounds in non-aqueous solvent, such as vegetable
- 10 oil, synthetic aliphatic acid glycerides, esters of
- 11 higher aliphatic acids or propylene glycol.

12

- 13 Formulations for topical use include known gels,
- 14 creams, oils, and the like. For aerosol delivery,
- 15 the compounds may be formulated with known aerosol
- 16 exipients, such as saline and administered using
- 17 commercially available nebulizers. Formulation in a
- 18 fatty acid source may be used to enhance
- 19 biocompatibility. Aerosol delivery is the preferred
- 20 method of delivery for epithelial airway
- 21 inflammation.

22

- 23 For rectal administration, the active ingredient may
- 24 be formulated into suppositories using bases which
- 25 are solid at room temperature and melt and dissolve
- 26 at body temperature. Commonly used bases include
- 27 cocoa butter, glycerinated gelatin, hydrogenated
- 28 vegetable oil, polyethylene glycols of various
- 29 molecular weights, and fatty esters of polyethylene
- 30 stearate.

- 1 The dosage form and amount can be readily
- 2 established by reference to known inflammatory
- 3 disease treatment or prophylactic regiments. The
- 4 amount of therapeutically active compound that is
- 5 administered and the dosage regimen for treating a
- 6 disease condition with the compounds and /or
- 7 compositions of this invention depends on a variety
- 8 of factors, including the age, weight, sex and
- 9 medical condition of the subject, the severity of
- 10 the disease, the route and frequency of
- 11 administration, and the particular compound
- 12 employed, the location of the inflammatory disease,
- 13 as well as the pharmacokinetic properties of the
- 14 individual treated, and thus may vary widely. The
- 15 dosage will generally be lower if the compounds are
- 16 administered locally rather than systemically, and
- 17 for prevention rather than for treatment. Such
- 18 treatments may be administered as often as necessary
- 19 and for the period of time judged necessary by the
- 20 treating physician. One of skill in the art will
- 21 appreciate that the dosage regime or therapeutically
- 22 effective amount of the inhibitor to be
- 23 administrated may need to be optimized for each
- 24 individual. The pharmaceutical compositions may
- 25 contain active ingredient in the range of about 0.1
- 26 to 2000mg, preferably in the range of about 0.5 to
- 27 500mg and most preferably between about 1 and 200
- 28 mg. A daily dose of about 0.01 to 100mg/kg body
- 29 weight, preferably between about 0.1 and about
- 30 50mg/kg body weight, may be appropriate. The daily
- 31 dose can be administered in one to four doses per
- 32 day.

- 2 Although the data presented is based predominantly 3 on the provision of cell lines that when induced
- 4 produce either homodimeric or heterodimeric IL-12,
- 5 or either subunit of IL-12, the invention is also
- 6 applicable in the production of cell lines which
- 7 when induced produce either IL-23 and IL-27, or
- 8 subunits thereof. In the case of IL-23, a suitable
- 9 host cell, such as one which includes an ecdysone-
- 10 inducible mammalian expression system as described
- 11 herein, is transformed with a first expression
- 12 vector according to the invention which includes DNA
- 13 coding for the p40 (beta) subunit of IL-12 (which is
- 14 identical to the p40 subunit of IL-23) and a second
- 15 expression vector which includes DNA coding for the
- 16 pl9 subunit of IL-23. In this regard, the cDNA
- 17 sequence of the p19 subunit of IL-23 is provided in
- 18 Sequence ID No. 8. The cDNA is processed by the same
- 19 restriction enzymes as used with the respective
- 20 subunits of IL-12, and is ligated into, for example,
- 21 a pIND vector is the same manner as is described
- 22 above. Likewise, expression vectors having DNA
- 23 coding for one of the subunits of Il-27, and cell
- 24 lines transfected with such expression vectors, may
- 25 be produced using the techniques described herein.

- 27 The invention is not limited to the embodiments
- 28 hereinbefore described which may be varied in detail
- 29 without departing from the invention.

1 Claims

2

- 3 1. An expression vector comprising DNA encoding a
- 4 subunit of a dimeric form of interleukin under
- 5 transcriptional control of an ecdysone-inducible
- 6 promoter.

7

- 8 2. A vector as claimed in Claim 1 in which the
- 9 subunit of a dimeric form of interleukin is selected
- 10 from the group comprising: p35 (alpha) subunit of
- 11 interleukin 12 (IL-12); p40 (beta) subunit of IL-12;
- 12 p19 chain of IL-23; p40 subunit of IL-23; ebi3
- 13 subunit of IL-27; and p28 subunit of Il-27.

14

- 15 3. A vector as claimed in Claim 1 or 2 comprising
- 16 an ecdysone-inducible mammalian expression plasmid,
- 17 wherein the DNA encoding the subunit of a dimeric
- 18 form of interleukin is included in the plasmid.

19

- 20 4. A vector as claimed in any preceding Claim in
- 21 which the DNA encodes a p40 subunit of IL-12.

22

- 23 5. A vector as claimed in any of Claims 1 to 3 in
- 24 which the DNA encodes a p35 subunit of IL-12.



- 1 6. A vector as claimed in any of Claims 1 to 3 in
- 2 which the DNA encodes a p19 subunit of IL-23.

- 4 7. An expression vector as claimed in Claim 1 or
- 5 6 in which the ecdysone inducible mammalian
- 6 expression vector is selected from the group
- 7 comprising: pIND; pIND(SP1); and pINDHygro.

8

- 9 8. An expression vector as claimed in any of
- 10 Claims 1 to 7 in which the DNA encoding a subunit of
- 11 dimeric interleukin 12 includes a DNA sequence
- 12 encoding a 6 x histidine tag.

13

- 14 9. An expression vector as claimed in any
- 15 preceding Claim selected from the group comprising:
- 16 pIND-p35H; pIND(SP1)-p35H; pIND-40H; pINDHygro-p40;
- 17 pIND(SP1)-p40H; and pIND-p40.

18

- 19 10. An expression vector as claimed in any
- 20 preceding Claim in which the DNA encoding the subunit
- 21 of dimeric interleukin is digested with NheI and XhoI
- 22 restriction enzymes prior to ligation of the digested
- 23 DNA products into the expression vector.

24

- 25 11. The expression vector pIND(SP1)-p35H having
- 26 ECACC accession number 03120401.

- 28 12. A method a producing a tightly controlled
- 29 expression vector capable of transforming a host cell
- 30 which when transformed is capable of producing a

- 1 recombinant dimeric interleukin, or a subunit
- 2 thereof, under transcriptional control of an
- 3 ecdysone-inducible promoter, comprising the steps of:
- 4 providing cDNA for a subunits of a dimeric
- 5 interleukin;
- 6 digesting the cDNA with at least one restriction
- 7 enzyme; and
- 8 ligating the digested cDNA product into an
- 9 ecdysone-inducible mammalian expression vector.

- 11 13. A method as claimed in Claim 12 in which the
- 12 one or more restriction enzymes consist of NheI and
- 13 XhoI.

14

- 15 14. A method as claimed in Claim 12 or 13 in which
- 16 the ecdysone-inducible mammalian expression vector is
- 17 selected from the group comprising: pIND; pIND(SP1);
- 18 and pINDHygro.

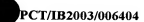
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- 20 15. A method as claimed in any of Claims 12 to 14
- 21 in which the cDNA for the subunit of dimeric
- 22 interleukin includes a DNA sequence encoding a 6  $\times$
- 23 histidine tag.

24

- 25 16. An expression vector obtainable by the method
- 26 of any of Claims 12 to 15.

- 28 17. A cell line transfected with at least one
- 29 expression vector of any of Claims 1 to 11 or 16,
- 30 wherein the DNA encoding the at least one subunit of



- 1 a dimeric interleukin is under the transcriptional
- 2 control of an ecdysone-inducible mammalian expression
- 3 system.

- 5 18. A cell line according to Claim 17 and capable
- 6 of producing homodimeric IL-12, the cell line being
- 7 transfected with an expression vector of Claim 4.

8

- 9 19. A cell line according to Claim 17 and capable.
- 10 of producing heterodimeric IL-12, the cell line being
- 11 transfected with an expression vector of Claim 4 and
- 12 an expression vector of Claim 5.

13

- 14 20. A cell line according to Claim 17 and capable
- of producing heterodimeric IL-23, the cell line being
- 16 transfected with an expression vector of Claim 4 and
- 17 an expression vector of Claim 6.

18

- 19 21. A cell line of any of Claims 17 to 20 which
- 20 includes a plasmid pVgRxR.

21

- 22 22. A cell line as claimed in any of Claims 17 to
- 23 21 in which the cells are human embryonic kidney
- 24 cells.

25

- 26 23. A cell line as claimed in Claim 22 in which
- 27 the cells are EcR293 cells.



- 1 24. A cell line as claimed in any of Claims 17 to
- 2 20 in which the cells are natural  $\beta$  subunit-producing
- 3 cells such as a HIBERNIA1 cell line.

- 5 25. A cell line having ECACC accession number
- 6 03112701.

7

- 8 26. A method of producing a cell line capable of
- 9 producing a recombinant dimeric interleukin, or a
- 10 subunit thereof, under transcriptional control of an
- 11 ecdysone-inducible promoter, comprising the steps of:
- 12 providing at least one expression vector
- according to any of Claims 1 to 11 or 16; and
- 14 transfecting a host cell with the at least one
- expression vector,
- 16 wherein the DNA encoding the at least one
- subunit of a dimeric interleukin is under the
- 18 transcriptional control of an ecdysone-inducible
- 19 mammalian expression system.

20

- 21 27. A method of preparing cDNA encoding a subunit
- 22 of a dimeric form of interleukin comprising the steps
- 23 of providing cDNA encoding the subunit, and digesting
- 24 the cDNA with restriction enzymes NheI and XhoI to
- 25 obtain a cDNA product.

- 27 28. A method of screening a candidate compound for
- 28 the ability to inhibit dimer assembly and secretion
- 29 of a dimeric form of interleukin, comprising the
- 30 steps of:



1 - incubating a cell culture comprising a cell line 2 of any of Claims 17 to 25 with the candidate 3 compound; 4 - inducing transcription of the dimeric 5 interleukin in the cells of the culture using 6 ecdysone or an ecdysone analog; and 7 - assaying the cell culture for the presence of 8 secreted interleukin. 9 A method as claimed in Claim 28, and in which 10 29. the interleukin expressed by the cell line has a 6  $\times$ 11 histidine amino acid sequence tagged on either or 12 13 both of the subunits thereof, wherein the assaying step involves Ni-NTA affinity chromatography. 14 15 A method as claimed in Claim 28 in which the 16 assaying step involves probing the cell culture with 17 18 an antibody specific to a dimeric form of interleukin, or a subunit thereof. 19 20 21 An inhibitor of dimer assembly and secretion of dimeric interleukin identified by the method of 22 23 any of Claims 28 to 30. 24 25 A method of prevention or treatment of inflammatory disease comprising a step of treating an 26 individual with an inhibitor of Claim 31. 27

33. A method of treating disease having apathogenesis which includes endogenous production of



- 1 any of cytokines IL-12, IL 23 or IL-27, the method
- 2 comprising a step of treating an individual with an
- 3 endoplasmic reticulum (ER) Ca<sup>2+</sup> perturbation reagent.

- 5 34. Use of an ER Ca<sup>2+</sup> perturbation reagent in the
- 6 manufacture of a medicament for the treatment of
- 7 disease having a pathogenesis which includes
- 8 endogenous production of any of cytokines IL-12, IL-
- 9 23 or IL-27.

10

- 11 35.. Use of an ER  $Ca^{2+}$  perturbation reagent for the
- 12 treatment of disease having a pathogenesis which
- 13 includes endogenous production of any of cytokines
- 14 IL-12, IL-23 or IL-27.

15

- 16 36. A method of inhibiting the formation of one or
- 17 more cytokines in an individual, which method
- 18 comprises the step of treating an individual with ER
- 19 Ca<sup>2+</sup> perturbation reagent.

20

- 21 37. Use of an ER Ca<sup>2+</sup> perturbation reagent to
- 22 inhibit the formation of one or more cytokines in an
- 23 individual.

24

- 25 38. A method or use as claimed in any of Claims 33
- 26 to 37 in which the disease is an inflammatory disease
- 27 in which one or more endogenously produced IL-12
- 28 forms play a disease promoting role.

- 1 39. A method or use as claimed in Claim 38 in
- 2 which the IL-12 forms are  $\alpha\beta$  heterodimeric and  $\beta\beta$
- 3 homodimeric forms.

4

- 5 40. A method or use as claimed in any of Claims 33
- 6 to 39 in which the disease is selected from the group
- 7 consisting of infectious diseases; bacterial
- 8 protozoal or virus-induced inflammation; epithelial
- 9 airway inflammation such as asthma; allergic disease;
- 10 autoimmune disease such as MS, RA and Inflammatory
- 11 Bowel Disease; and -all conditions in which
- 12 endogenously produced IL-12  $\alpha/\beta$  or  $\beta\beta$  forms are
- 13 thought to play a disease-promoting role.

14

- 15 41. A method or use as claimed in any of Claims 33
- 16 to 40 in which the ER  $Ca^{2+}$  perturbation reagent is
- 17 selected from the compounds of Formula I:

18

19

20 Formula I

$$R^2$$
  $S^{O}$   $R^3$ 

21

- 22 wherein A is a substituent selected from partially
- 23 unsaturated or unsaturated hetrocyclyl and partially
- 24 unsaturated or unsaturated carbocyclic rings;
- 25 wherein  $R^1$  is at least one substituent selected from
- 26 hetercyclyl, cycloalkyl, cycloalkenyl and aryl,

- 1 wherein R1 is optionally substituted at a
- 2 substitutable position with one or more radicals
- 3 selected from alkyl, haloalkyl, cyano, carboxyl,
- 4 alkoxycarbonyl, hydroxyl, hydroxyalkyl, amino,
- 5 alkylamino, arylamino, nitro, alkoxyalkyl,
- 6 alkylsulfinyl, halo, alkoxy and alkylthio;
- 7 wherein  $R^2$  is methyl or amino; and
- 8 wherein R<sup>3</sup> is a radical selected from hydrido, halo,
- 9 alkyl, alkenyl, oxo, cyano, carboxyl, cyanoalkyl, .
- 10 heterocyclyloxy, alkyloxy, alkylthio, alkylcarbonyl,
- 11 cycloalkyl, aryl, haloalkyl, heterocyclyl,
- 12 cycloalkenyl, aralkyl, hetrocyclylalkyl, acyl,
- 13 alkythioalkyl, hydroxyalkyl, alkoxycarbonyl,
- 14 arylcarbonyl, aralkylcarbonyl, aralkenyl,
- 15 alkoxyalkyl, arylthioalky, aryloxyalkyl,
- 16 aralkylthioalky, aralkoxyalkyl, alkoxyaralkoxyalkyl,
- 17 alkoxycarbonalkyl, aminocarbonyl, aminocarbonylalkyl,
- 18 alkyaminocarbonyl, N-arylaminocarbonyl, N-alkyl-N-
- 19 arylaminocarbonyl, alkylaminocarbonylalkyl,
- 20 carboxyalkyl, alkylamino, N-arylamino, N-
- 21 aralkylamino, N-alkyl-N-aralkylamino, N-alkyl-N-
- 22 arylamino, aminoalkly, alkylaminoalkyl, N-
- 23 arylaminoalkyl, N-aralkylaminoalkyl, N-alkyl-N-
- 24 aralkylaminoalky, N-alkyl-N-arylaminoalkyl, aryloxy,
- 25 aralkoxy, arylthio, aralkylthio, alkylsulfinyl,
- 26 alkylsulfonyl, aminosulfonyl, alkylaminosulfonyl, N-
- 27 arylaminosulfonyl, arylsulfonyl, N-alkyl-N-
- 28 arylaminosulfonyl; or a pharmaceutically-acceptable
- 29 salt thereof.

1/9

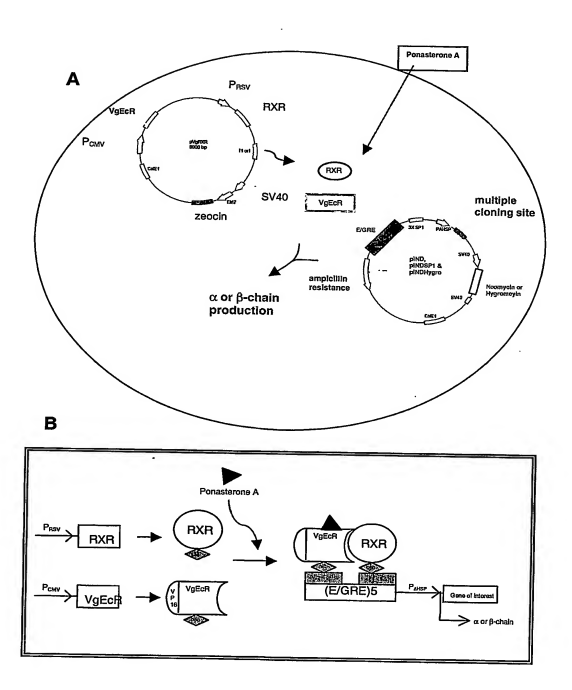


Fig. 1.

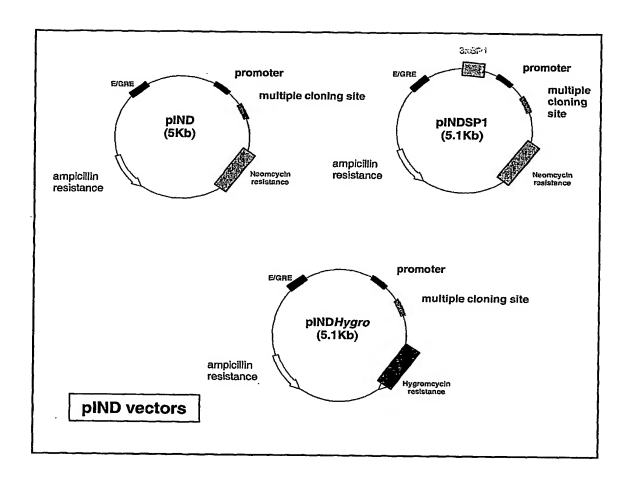


Fig. 2.

3/9

- A 5' CAGGCT AGG GCA GCC ATG TGT CCA GCG CGC AGC 3'
- B 5' CTGCTC GAG TTA ATG GTG ATG GTG ATG GTG GGA AGC ATT CAG ATA GCT 3'

  Xhol restriction site

Nhel, restriction site

- C 5' CAG GCT AGG GCA GCC ATG TGT CAC CAG CAG TTG
- D 5' CTGCTC GAG CTA ATG GTG ATG GTG ATG GTG ACT GCA GGG CAC AGA TG 3'

  Xhol restriction site
- E 5' CTG CTC GAG CTA ACT GCA GGG CAC AGA TG 3'

  Xhol restriction site

Fig. 3.

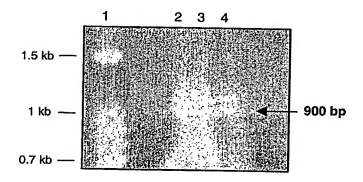


Fig. 4.

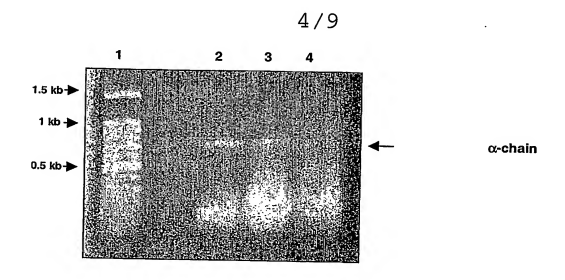


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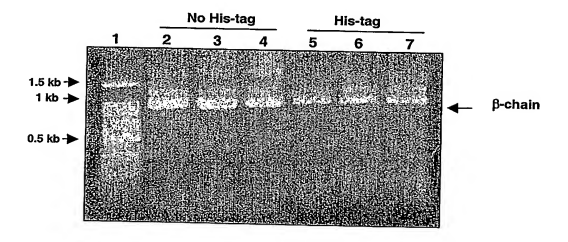
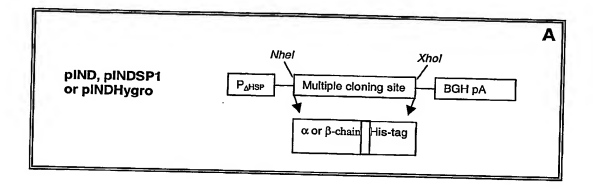
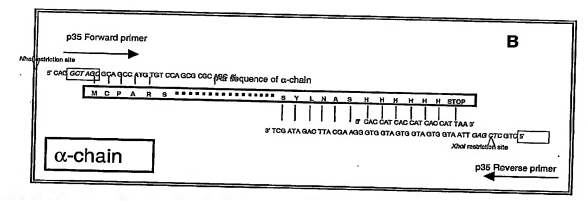


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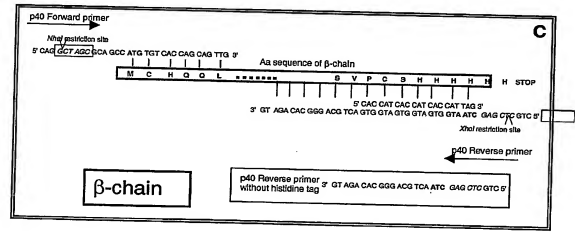
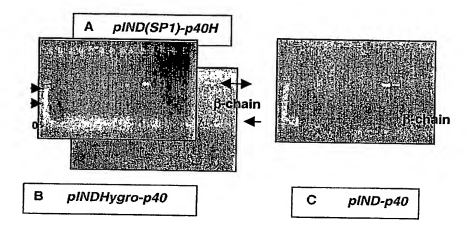


Fig. 7.

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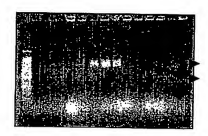


Fig. 9.

**ፈ**\_α-chain

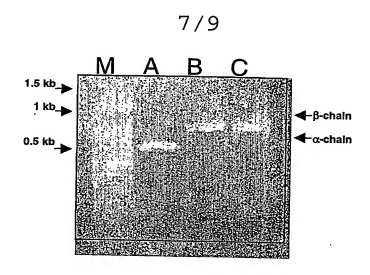


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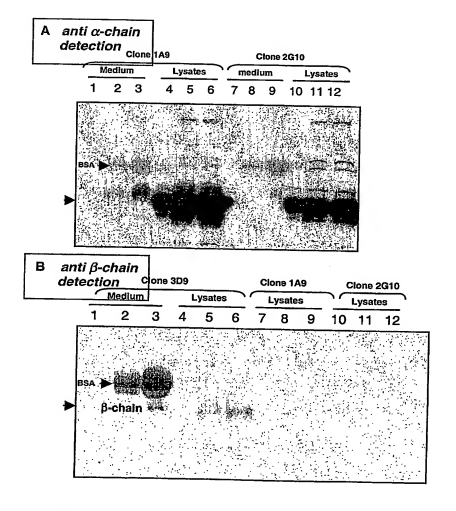


Fig. 11

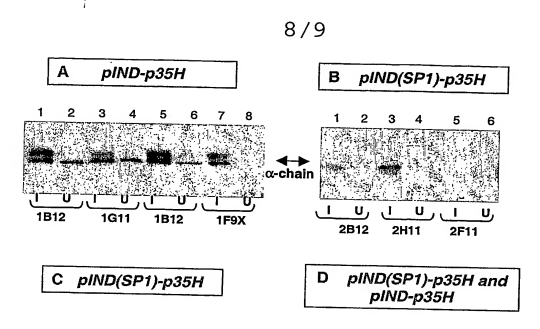


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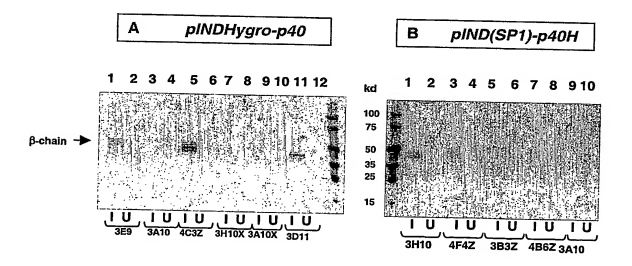


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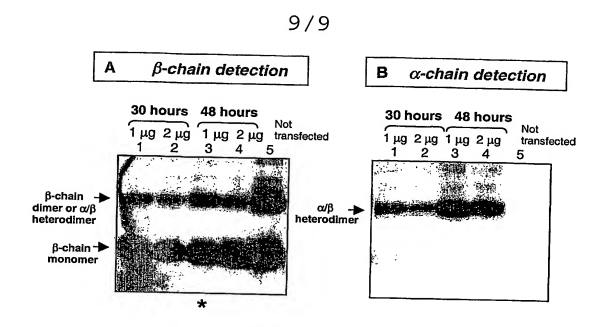


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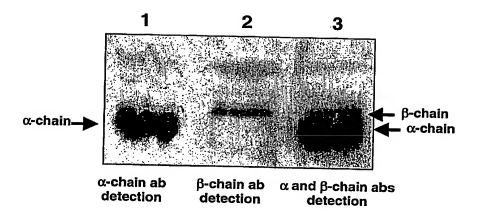


Fig. 15.

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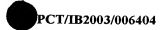
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